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cDNA; 713 BP. secreted and transmembrane protein PRO1159
          Length 713;
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                                           ACA64409 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
                                                                                                                                               Length 713;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human secreted/transmembrane protein (PRO) cDNA #237.
US2003036179-A1.
                                                                                                                                                                                                                                          20-25-2003.
20-25-2003.
(GETH) GENENTECH INC.
100.0%; Score 273; DB 8;
tery Match 100.0%; Pred. No. 3.3e-75;
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Human sectreted/transmembrane protein cDNA, #154
US2002142961-A1.
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Pred. No. 3.3e-75;
          100.0%; Score 273; DB 5; 100.0%; Pred. No. 3.3e-75;
                                                                                                                                               273; DB 8;
No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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No. 3.3e-75;
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Human secreted/transmembrane protein cDNA, #154
US2003027162-A1.
06-FEB-2003.
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Pred. No. 3.3e-75
                                                                                                                                                                                                                 #237
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cDNA encoding human PRO1159 polypeptide
US2002127576-A1.
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Pred.
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Pred.
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(GETH ) GENENTECH INC.
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02-JAN-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                    ABX89377 standard;
DNA encoding novel
US2003017563-A1.
                       Best Local Similarity RESULT 6
                                                                                                                                               Query Match
Best Local Similarity
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GenCore version 5.1.6
(c) 1993 - 2004 Compugen Ltd.
2004, 00:55:08; Search time 204.612 Seconds
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7003.964 Million cell updates/sec
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SUMMARIES
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Human PRO1159 protein UNQ589 encoding cDNA SEQ ID NO:272
WO200053758-A2.
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Human PRO1159 (UNQ589) nucleotide sequence SEQ ID NO:376
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Human cDNA sequence encoding for PRO1159 polypeptide.
WO200140466-A2.
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Membrane-bound protein PRO1159 encoding cDNA
WO9963088-A2.
                                                                                         US-09-989-293A-376_COPY 92 364
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14-SEP-2000.
(GETH ) GENENTECH INC.
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December 2,
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 4
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Best Local Similarity
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ACA04260 standard; cDNA; 713

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Human cDNA encoding a secreted/transmembrane protein, SEQ ID 473
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                                                                       Length 713
                                                                                                                                                                                                                                                                                                                                                                                                 ACA68006 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2002177164-A1.
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                                                                                                                            ACA93067 standard, cDNA, 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003017476-A1.
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Human secreted and transmembrane polypeptide PRO1159 cDNA
US2002197615-A1.
                                                                       100.0%; Score 273; DB 8; 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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cDNA encoding human PRO1159 polypeptide.
US2003017981-A1.
                                                                                                                                                                                                                                                      ABX17151 standard; cDNA; 713 BP. Human PRO polynucleotide #118. US2002123463-A1.
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W22003049816-A1.
13-MAR-2003.
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Human PRO polynucleotide #237.
US2003073212-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC
                                                        (GETH ) GENENTECH INC
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Best Local Similarity
RESULT 21
ID ADA45992 standard, cl
DE Novel human secreted
PN UG200302238-A1.
PD 30-JAN-2003.
PA (GETH ) GENENTECH INC
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                                                                                    Best Local Similarity RESULT 16
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                US2003032155-A1.
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ID ACA88
DE Human
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PD 26-DE
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Human cDNA encoding secreted/transmembrane polypeptide PRO1159. 20-MAR-2003.
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Human cDNA encoding secreted/transmembrane protein, PRO1159
US200305981-A1.
27-MAR-2003.
                                                                                   ADB19481 standard; cDNA; 713 BP. Novel human secreted and transmembrane protein PRO1159 cDNA
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082711-A1.
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                                                                                              Novel human ...
US2003068796-A1.
US-003068796-A1.
(GETH ) GENENTECH INC.

"a+ch 'l~vity 100.0%; Score 273; DB 9;
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(GETH ) GENENTECH INC.
iry Match
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(GETH ) GENENTECH INC.
ery Match 100.0%; Score 273; DB 9;
st Local Similarity 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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US2003082704-A1.
01-MAY-2003.
(EETH ) GENENTECH INC.
ERY MACCh 100.0%; Score 273;
Bt Local Similarity 100.0%; Pred. No. 3
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Human PRO polynucleotide #237.
US2003087350-A1.
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US2003073215-A1.
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17-APR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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                              Query Match
Best Local Similarity
RESULT 25
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Best Local Similarity
RESULT 31
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Best Local Similarity
RESULT 32
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Best Local Si
RESULT 28
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Query Match

RESULT

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Length 713;
                                                                     Novel human secreted and transmembrane protein PRO1159 cDNA. 17-APR-2003. (Appr. 2003).
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082691-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003044945-A1.
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Muman cDNA encoding secreted/transmembrane protein PRO1159. US2003059812-A1.
27-MAR-2003.
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(GETH ) GENENTECH INC.
(GETY Match 100.0%; Score 273; DB 9; Lery Match 100.0%; Pred. No. 3.3e-75;
                                                100.0%; Score 273; DB 9; 100.0%; Pred. No. 3.3e-75;
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100.0%; Score 273; DB 9;

17 Match

17 Acal Similarity 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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ID NO 473
                                                                                                                                                                                                                                      Human PRO polynucleotide #237.
24-APP. A..
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06-MAR-2003.
(GETH ) GENENTECH INC.
100.0%;
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ID ADB24832 standard, cDNA, 713
DE Human PRO polymucleotide SEQ
PN US2003077713-A1.
             08-MAY-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
                                                             Best Local Similarity RESULT 44
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Best Local Similarity
RESULT 49
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Best Local Similarity
RESULT 50
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US2003087351-A1
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PN
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Human PRO polynucleotide #237.

US2003082763-A1.

01-MAY-2003.

(GETH ) GENENTECH INC.

ery Match

st Local Similarity 100.0%; Pred. No. 3.3e-75;
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Novel human secreted and transmembrane protein PRO1159 cDNA, US2003087145-A1.
(88-MAY-2003.
(GETH ) GENENTECH INC.
                                                                                                       Length 713
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Novel human secreted and transmembrane protein PRO1159 cDNA US2003082694-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1159
US2003054359-A1.
                                                                                                                                                                        Novel human secreted and transmembrane protein PRO1159 US2003082693-A1.
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(GETH ) GENENTECH INC.

(GETH ) GENENTECH INC.

ELY Match 100.0%; Score 273; DB 9;

Cimilarity 100.0%; Pred. No. 3.3e-75;
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01-MAY-2003.
(GETH ) GENENTECH INC.
(GETH ) SCORE 273; DB 9;
ery Match
' rimilarity 100.0%; Pred. No. 3.3e-75;
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01-MAY 2003.
(GETH ) GENENTECH INC.
100.0%; Score 273; DB 9;
LETY MATCh 100.0%; Pred. No. 3.3e-75;
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08-MAY-2003.

(GFH ) GENENTECH INC.

(GFT ) SCORE 273; DB 9;

GFT MATCH 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
           ADB30653 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237. US2003068794-A1.
                                                                                                                                                                                                                                                                                                  ADA17904 standard; cDNA; 713 BP. cDNA encoding human PRO1159 polypeptide. V22003054987-A1.
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Human PRO polynucleotide #237.
                                                                                                                                                         CDNA; 713 BP
                                                                                  INC.
100.0%;
Y 100.0%;
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RESULT 39
10 ADA87604 standard; CL
DE Novel human secreted
PN US200308745-A1.
PD 08-MAY-2003
PA (GETH ) GENENTECH INC
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Query Match
Best Local Similarity
RESULT 42
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Best Local Similarity
RESULT 36
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Best Local Similarity
RESULT 37
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Best Local Similarity
RESULT 38
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Best Local Similarity
RESULT 41
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Best Local Similarity
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                                                             10-APR-2003.
(GETH ) GENENTECH
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Query Match

Query Match

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Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA US2003082695-A1.
01-MAY-2003.
                                                                                                                                                                                                                                                                              Length 713
                                                                                                                                                                                                                                                                                                                         ADA@4845 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082708-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1159. US2003059780-A1.
27-MAR-2003.
(GETH ) GENENTECH INC.
100.0%; Score 273; DB 9;
100.0%; Pred. No. 3.3e-75;
                                                                                                               17-APR-2003.
(GETH) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 9;
ery Match 100.0%; Pred. No. 3.3e-75;
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(GETH) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 9;
ery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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282003073214-Al.
17-APR-2003.
(GETH ) GENENTECH INC.
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Human PRO polynucleotide SEQ ID NO 473.
US2003077715-Al.
                                                                    ADA75319 standard, cDNA; 713 BP. Human PRO polynucleotide #237. US2003073216-A1.
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Human PRO polynucleotide #237.
US2003082703-A1.
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PA (GETH ) GENENTECH INC.

Query Match 100.0%; Sc
Best Local Similarity 100.0%; Px
RESULT 56

ID ADB30101 standard; CDNA; 713 BP.
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US2003082761-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 61
ID ADB25392 standard; cl
DE Human PRO polymucleot
PD US2003077715-A1.
PD 24-APR-2003.
PA (GETH ) GENENTECH INC
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Best Local Similarity
RESULT 58
                        Query Match
Best Local Similarity
RESULT 53
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RESULT 54
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Length 713;
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Muman cDNA encoding secreted/transmembrane protein PRO1159. US200306407-A1.
27-MAR-2003.
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01-MAY-2003.

(GFTH) GENENTECH INC.

(GFTH) GENENTECH INC.

100.0%; SCORE 273; DB 9;

100.0%; Pred. No. 3.3e-75;
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Best Local Similarity 100.0%; Pred. No. 3.3e-75; RESULT 62
                                                                                                                                                                                                                                                    Score 273; DB 9;
Pred. No. 3.3e-75;
                                                                                                                     Score 273; DB 9;
Pred. No. 3.3e-75;
                                                                                                                                                                                                                                                                                                                 cDNA encoding human PRO polypeptide #237.
US2003096386-A1.
22-MAY-2003.
(GETH ) GENENTECH INC.
100.0%; Score 273; DB 9; Bt Local Similarity 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
                                                                                                                                                                                   cDNA encoding human PRO polypeptide #237 US2003092147-A1.
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Human PRO polynucleotide SEQ ID NO 473.
US200307714-A1.
24-ARR-2003.
(GETH ) GENENTECH INC.
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01-MW- ....
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Human PRO polynucleotide #237.
US2003082690-A1.
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Human PRO polynucleotide #237.
US2003082759-A1.
                                     ADA93568 standard; cDNA; 713 BP.
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24-APR-2003.
(GETH ) GENENTECH INC.
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01-MAY-2003.
(GETH ) GENENTECH INC.
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                                                 Human PRO polynucleotide #237.
US2003077721-A1.
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15-MAY-2003.
(GFH) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                                     Query Match
Best Local Similarity
RESULT 63
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RESULT 68
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Best Local Similarity
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US2003049817-A1.
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RESULT 67
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ADA22499 standard; cDNA; 713 BP.
Human cDNA encoding secreted/transmembrane polypeptide PRO1159.
US2003040473-A1.
27-FEB-2003.
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   encoding secreted/transmembrane protein PR01159
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Novel human secreted and transmembrane protein PRO1159 US2003073213-A1.
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(GETH ) GENENTECH INC.
iry Match 100.0%; Score 273; DB 9;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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                                                                               Human PRO polynucleotide #237.
US2003059909-A1.
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(GETH) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                         Best Local Similarity RESULT 81
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Human cDNA encodus2003044806-A1
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(GETH ) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 9; Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003054516-A1.
20-MAR-2003.
YOUR GENENTECH INC.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082765-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082709-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082700-A1.
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(GETH) GENENTECH INC.
ery Match 100.0%; Score 273; DB 9;
ery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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No. 3.3e-75;
 ADB26366 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2003082760-Al.
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Pred.
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US2003068797-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 80
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Best Local Similarity
RESULT 79
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Best Local Similarity
RESULT 75
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RESULT 76
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Best Local Similarity
RESULT 72
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                                               01-MAY-2003
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RESULT 78

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Length 713;

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PA (GETH) GENENTECH INC.

Query Match 100.0%; Score 273; DB 10; Length 713;

Best Local Similarity 100.0%; Pred. No. 3.3e-75;

RESULT 100
                                                                                                                                                                     Score 273; DB 10; Length 713; Pred. No. 3.3e-75;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082764-A1.
                                                                                                                                                                                                                                                                                                                                       AUD4/221 standard; CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA
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Novel human secreted and transmembrane protein PRO1159 US2003082696-A1.
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(GETH) GENENTECH INC.
(ery Match
'cery match' of milarity 100.0%; Pred. No. 3.3e-75;
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24-APR-2003.
(GETH ) GENENTECH INC.
100.0%; Score 273; DB 10;
10T Match 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.

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(ery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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ID NO 473.
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Human PRO polynucleotide SEQ ID NO 473.
US2003077719-A1.
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Human PRO polynucleotide SEQ ID NO 473.
US2003077716-A1.
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Human PRO polynucleotide SEQ ID NO 473.
US2003077718-A1.
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Human PRO polynuclectide #237.
US20082697-A1.
01-MAY-2003.
GETH ) GENENTECH INC.
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est Local Similarity 100.0%; Scor
                                                                           Human PRO polynucleotide #237.
01-MAY-2001
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24-APR-2003.
(GETH ) GENENTECH INC.
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01-MAY-2003.
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Human PRO polynucleotide
US2003077717-A1.
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(GETH ) GENENTECH INC.
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RESULT 102
ID ADB47221 standard;
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RESULT 106
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(GETH ) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 9; Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082712-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082766-A1.
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Novel human secreted and transmembrane protein PRO1159 CDNA
US2003082689-A1.
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Novel human secreted and transmembrane protein PRO1159
US2003087347-A1.
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(GETH) GENENTECH INC.

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                                                                 100.0%; Score 273; DB 9; 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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                                                                                                                                                                                                                                               ADB23728 standard; cDNA; 713 BP.
Human PRO polynucleotide SEQ ID NO 473.
US2003077712-A1.
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Human PRO polynucleotide #237.
US2003087352-A1.
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24-APR-2003.
(GETH ) GENENTECH INC.
100.0%; SC
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Human PRO polynucleotide #237.
US2003077711-A1.
Human PRO polynucleotide #237.
US2003066793-A1.
10-APR-2003
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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ID ADB15513 stand*

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Best Local Similarity
RESULT 99
ID ADB89765 Streep
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RESULT 97
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Best Local Similarity
RESULT 91
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Best Local Similarity
RESULT 93
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ADC53047 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein cDNA Seg ID473
US203087355-A1.
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ID ADC54692 standard; CDNA; 713 BP.
DE Novel human secreted and transmembrane protein CDNA Seq ID473 PN US2003087363-A1.
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Novel human secreted and transmembrane protein cDNA Seg ID473.
US2003087364-A1.
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Novel human secreted and transmembrane protein cDNA Seq ID473
US2003087359-A1.
08-MAY-2003.
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Novel human secreted and transmembrane protein PR01159 cDNA.
US2003087367-A1.
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Novel human secreted and transmembrane protein cDNA Seq
US2003087366-A1.

08-MAY-2003.
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                                                                                     ADC60040 standard; cDNA; 713 BP. Novel human secreted and transmembrane protein PRO1159 US2003092105-A1.
15-MAY-2003.
(GETH ) GENENTECH INC.
100.0%; Score 273; DB 10;
ATT 100.0%; Pred. No. 3.3e-75;
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No. 3.3e-75;
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Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
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RESULT 126
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RESULT 119
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RESULT 120
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Best Local Similarity
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ID ADC65594 standard;
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RESULT 121
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082692-A1.
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RESULT 116
ID ADC11554 standard; CDNA; 713 BP.
DE Human cDNA encoding secreted/transmembrane protein PRO1159.
PN US2003069403-A1.
PD 10-APR-2003.
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Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003049681-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1159.
US200306847-A1.
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(GETH ) GENENTECH INC.
sry Match 100.0%; Score 273; DB 10;
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Pred. No. 3.3e-75;
                                                                         ADB36246 standard; cDNA; 713 BP.
Human PRO polynucleotide SEQ ID NO 473.
US2003077720-A1.
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D ADG57856 standard; CDNA; 713 BP. DE Human PRO polynucleotide #118.
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Human PRO polynucleotide #118.
US2003064375-A1.
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(GETH ) GENENTECH INC.
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RESULT 118
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Best Local Similarity
RESULT 110
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RESULT 109
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RESULT 112
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Query Match

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ADD10127 standard, cDNA, 713 BP.
Human PRO polynucleotide #237.
US2003194776-A1.
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RESULT 145
  DE DE LO
                                                   ADC56054 standard; cDNA, 713 BP.

Novel human secreted and transmembrane protein cDNA Seq ID473.
US2003087360-A1.
08-MAY-2003.
(GETH ) GENENTECH INC.
100.0%; Score 273; DB 10; Length 713; St Local Similarity 100.0%; Pred. No. 3.3e-75;
                                                                                                                                                                   ADCS8624 standard; cDNA; 713 BP.

Novel human secreted and transmembrane protein cDNA Seq ID473
US2003087346-A1.
              Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003068623-A1.
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Novel human secreted and transmembrane protein PR01159 cDNA.
US2003092104-A1.
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Novel human secreted and transmembrane protein PR01159
US2003087348-A1.
             Score 273; DB 10;
Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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No. 3.3e-75;
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14-OCT-2003.
(GETH ) GENENTECH INC.
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cDNA encoding human PRO polypeptide #237.
US2003194770-A1.
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Pred.
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Pred.
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Human PRO polynucleotide #118.
US2003083461-A1.
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Human PRO polynucleotide #237.
US2003194773-Al.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 134
                      Best Local Similarity
RESULT 128
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Best Local Similarity
RESULT 130
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Best Local Similarity
RESULT 137
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RESULT 131
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Best Local Similarity
RESULT 135
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                                                                                                                                                                                                   ADD07675 standard, cDNA, 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2002193299-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003087134-Al.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003092103-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003073090-A1.
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Query Match
Best Local Similarity 100.0%; Pred. No. 3.3e-75;
RESULT 143
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(GBTH ) GENENTECH INC.
(GBTH ) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 10;
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Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
ery Match 100.0%; Score 273; DB 10;
ery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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27-MAD. Action | Hiller | 
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US2003194771-A1.
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SACCAL
16-OCT-2003.
(GETH) GENENTECH INC.
"Match "T--itv 100.0%;
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US-cv-c-
(GETH ) GENENTECH INC.
100.0%;
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Best Local Similarity 100.0%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 140
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RESULT 141
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Length 713;

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19-SEP-2002.
(GETH ) GENENTECH INC.
ery Match 100.0%; Score 273; DB 10; Length 713;
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Best Local Similarity 100.0%; Pred. No. 3.3e-75;
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ID ADE32414 standard; cDNA; 713 BP.
DE Novel human secreted and transmembrane protein PRO1159 cDNA.
PN US2003194765-A1.
                                                                                                                                                                                                                                                      Best Local Similaria,
RESULT 158

1D ADD5290 standard, cDNA, 713 BP.
DE Novel human secreted and transmembrane protein PRO1159
PN US2003203432-A1.
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(GETH ) GENENTECH INC.
2ry Match
2ry Match 100.0%; Score 273; DB 10;
2ry Match 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
Lery Match 100.0%; Score 273; DB 10;
Lery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 10;
ery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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                                                                                                                                          Human PRO polynucleotide #237.
US2003203430-A1.
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RESULT 156

ID ADD02674 standard; cDNA; 713 BP.

DE Human PRO polynucleotide #237.

PN US2003203431-A1.
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ID ADD54745 standard; CDNA; 713 BP.
DE Human PRO polynucleotide #118.
PN US2002132233-A1.
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Human PRO polynucleotide #237.
US2003199057-A1.
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30-0CT-2003.
(GETH ) GENENTECH INC.
"atch "letty 100.0%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                               Best Local Similarity
RESULT 157
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ID ADD92607 standard;
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ID ADE26899 standard;
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                                                                                                                                                                                                                                                                                                                                              ADD41288 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003203438-A1.
30-OCT-2003.
(GETH) GENENTECH INC.
100.0%; Score 273; DB 10; Length 71:
st Local Similarity 100.0%; Pred. No. 3.3e-75;
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ID ADD53119 standard; cDNA; 713 BP.
DE Novel human secreted and transmembrane protein PRO1159
PN US2003203437-A1.
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US2003194779-Al.
16-OCT-2003.
(GETH ) GENENTECH INC.
ery Match
st Local Similarity 100.0%; Pred. No. 3.3e-75;
   PD 19-DEC-2002.

PA (GETH ) GENENTECH INC.

Query Match

100.0%; Score 273; DB 10;

Best Local Similarity 100.0%; Pred. No. 3.3e-75;

RESULT 147
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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cDNA encoding human PRO polypeptide #237.
US2003194769-A1.
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cDNA encoding human PRO polypeptide #237.
US2003194792-A1.
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                                                                                  ADD09575 standard, cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003194775-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC
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Best Local Similarity
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Best Local Similarity
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RESULT 149
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Best Local Similarity
RESULT 151
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Best Local Similarity
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RESULT 153
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RESULT 154
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Length 713;

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PA (GETH ) GENENTECH INC.

QUETY MATCH

Best Local Similarity 100.0%; Score 273; DB 10; Length 713;

RESULT 176
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                23-0CT-2003.
(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
ery Match 100.0%; Score 273; DB 10;
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23-OCT_2003.
(GETH ) GENENTECH INC.
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100.0%; Score 273; DB 10;
100.0%; Pred. No. 3.3e-75;
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PA (GETH ) GENENTECH INC.

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100.0%; Score 273; DB 10;

RESULT 182.

RESULT 182.
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Human PRO polynucleotide #237.
US2001199059-A1.
23-OCT-2003.
(GETH ) GENENTECH INC.
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EY WALCH.
EY LOCAl Similarity 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
Lery Match 100.0%; Score 273; DB 10;
Lery Match 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC,
lery Match
100.0%; Score 273; DB 10;
100.0%; Pred. No. 3.3e-75;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADE22898 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237. US2003199064-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADD79016 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237. US2003203429-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADD80674 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237.052003207418-A1.
                                                                                                                         ADE19027 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003199026-A1.
                                                                                                                                                                                                                                                                                 ADE41223 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003199033-A1. 23-CCT-2003. (GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADE42658 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003199032-A1.
                                                                                 Best Local Similarity RESULT 175
                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 177
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Best Local Similarity
RESULT 178
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      US2003199025-A1.
                                                                    Query Match
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RESULT 179
                                                                                 Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003194767-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA, US2003194791-A1.
16-OCT--2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                            100.0%; Score 273; DB 10; 100.0%; Pred. No. 3.3e-75;
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Human PRO polynucleotide #237.
22-0CT-2003.

(GETH) GENENTECH INC.

ery Match

st Local Similarity 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
12ry Match 100.0%; Score 273; DB 10;
1. 700al Similarity 100.0%; Pred. No. 3.3e-75;
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PA (GETH ) GENENTECH INC.

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Best Local Similarity 100.0%; Pred. No. 3.3e-75;

RESULT 168
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                                                                           Query Match
Best Local Similarity 100.0%; Score 273;
Best Local Similarity 100.0%; Pred. No. 3
TD AD19570 standard; CDNA; 713 BP.
DB CDNA encoding human PRO polypeptide #237.
PN US20032032428-A1.
PA (GETH ) GENENTECH INC.
cDNA encoding human PRO polypeptide #237.
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                                                                                                                                                                                                                                                                                                  ADE42106 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003194772-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADD92055 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2001199053-A1. 23-OCT-2003.
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US2003194768-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 171
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RESULT 170
                  US2003199056-A1.
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ID ADD931
DE Human
PN US2003
PD 16-OCT
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100.0%; Score 273; DB 10; Length 713; 100.0%; Pred. No. 3.3e-75;
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                                                                                                                                                                                    ADH55403 standard; cDNA; 713 BP.

Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207381-A1.
06-NOV-2003.
(GETH) GENENTECH INC.
100.0%; Score 273; DB 10; Length 713
st Local Similarity 100.0%; Pred. No. 3.3e-75;
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207386-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2002207387-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207388-A1.
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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(GETH) GENENTECH INC.
iry Match 100.0%; Score 273; DB 10;
'' ramal Similarity 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Human PRO polynucleotide #118.
US2003050457-A1.
                                                   ADG80111 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003207372-A1.
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06-NOV-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 200
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   Query Match
Best Local Similarity
RESULT 193
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Best Local Similarity
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RESULT 196
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ID ADI63622 standard;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207384-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADG21623 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207355-A1.
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(GENNY-2003.

(GETH) GENENTECH INC.

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100.0%; Score 273; DB 10;

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100.0%; Pred. No. 3.3e-75;
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23-OCT-2003.
(GETH ) GENENTECH INC.
ery Match
100.0%; Score 273; DB 10;
ery Match
100.0%; Pred. No. 3.3e-75;
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Human PRO polynucleotide #237.

105203199031-A1.

23 -OCT-2003.

(GETH) GENENTECH INC.

100.0%; Score 273; DB 10; st Local Similarity 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
:ry Match 100.0%; Score 273; DB 10;
:ry Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
                     100.0%; Score 273; DB 10; 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Human PRO1159 nucleotide sequence SEQ ID NO:376.
US2002198148-A1.
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                                                                                                                                                                                                                                                                                                                                 Human PRO polymucleotide #237. US2003199034-A1.
                                                                     ADD89702 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003199028-A1.
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06-NOV-2003.
(GETH) GENENTECH INC.
Match 'Towity 100.0%;
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16-OCT-2003.
(GETH ) GENENTECH INC.
100.0%;
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26-DEC-2002.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC
       (GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 191
ID ADF97599 standard; c:
DE Human PRO polynucleo
PN US2003207370-A1.
PD 06-NOV-2003.
PA (GETH ) GENENTECH IN
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Best Local Similarity
RESULT 185
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Best Local Similarity
RESULT 189
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Best Local Similarity
RESULT 190
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Best Local Similarity
RESULT 192
                     Query Match
Best Local Similarity
RESULT 184
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RESULT 187
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Best Local Similarity
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                                                                                           ADNIE681 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003087385-A1.
                                                                                                                                                                                                                          ADN15500 standard; CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA.
US2003087356-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADC81210 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003022115-A1.
15-MAY-2003.
(GFTH) GENENTECH INC.
ery Match
st Local Similarity 100.0%; Pred. No. 3.3e-75;
                                                                                                                                                                                                                                                                                                                                                                             ADM14948 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 US2003087357-A1.
     PD 08-MAY-2003.

PA (GETH ) GENENTECH INC.

Query Match 100.0%; Score 273; DB 11;
Best Local Similarity 100.0%; Pred. No. 3.3e-75;
RESULT 212
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(GETH ) GENENTECH INC.
Hery Match 100.0%; Score 273; DB 12;
ery Match 100.0%; Pred. No. 3.3e-75;
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(GETH) GENENTECH INC.
(GETY MATCh 100.0%; Score 273; DB 11;
ery Match 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
100.0$; Score 273; DB 12;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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CDNA encoding human PRO polypeptide #237.
US2003092108-Al.
15-MAY-2003.
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Human PRO polynucleotide #237.
US2003092113-A1.
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08-MAY-2003.
(GETH ) GENENTECH INC.
100.0%;
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13-WOV-2003.
(GETH ) GENENTECH INC.
100.0%;
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(GETH ) GENENTECH INC.
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RESULT 216
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                                                                                                                                                                                                                      Score 273; DB 10; Length 713; Pred. No. 3.3e-75;
                                                                                           100.0%; Score 273; DB 10; Length 713; 100.0%; Pred. No. 3.3e-75;
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ADH81484 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US200320737-A1.
(GENV-2003.
(GETH ) GENENTECH INC.
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Novel human secreted and transmembrane protein PRO1159 CDNA.
US2003032156-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003087355-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003087353-A1.
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Wham cDNA encoding secreted/transmembrane protein PRO1159.
US2003032023-A1.
13-PEB-2003.
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Novel human secreted or transmembrane protein PRO1124 DNA
US2002132252-A1.
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(GETH ) GENENTECH INC.
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.... 100.0%; Score 273; DB 10;
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(GETH ) GENENTECH INC.

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100.0%; Score 273; DB 10;

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100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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protein cDNA, #154
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cDNA encoding human PRO polypeptide #237.
US2003004311-A1.
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CDNA encoding human PRO1159 polypeptide.
US2002103125-A1.
                                                                                                                                    Human PRO polynucleotide #118. US2003027163-A1.
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ID ABX90341 standard; CDNA; 713 BP DE Human secreted/transmembrane pr PN US2002160384-A1.
PD 31-CCT-2002.
PA (GETH ) GENENTECH INC.
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19-SEP-2002.
(GETH ) GENENTECH INC.
100.0%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 209
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RESULT 205
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RESULT 206
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RESULT 203
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100.0%; Pred. No. 3.3e-75;
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Best Local Similarity RESULT 230
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Best Local Similarity
RESULT 231
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Best Local Similarity
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RESULT 234
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Best Local Similarity
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(GETH ) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 12;
(ery Match 100.0%; Pred. No. 3.3e-75;
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15-MAY-2003.
(GETH) GENENTECH INC.
(GETY MATCh
(Ery Match 100.0%; Score 273; DB 12;
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
ery Match
' rimilarity 100.0%; Pred. No. 3.3e-75;
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(GETH) GENENTECH INC.
100.0%; Score 273; DB 12;
ery Match
100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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                    Score 273; DB 12;
Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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cDNA encoding human PRO polypeptide #237.
US2003199027-A1.
                                                                                                                                                                               ADE24645 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237.US2003092111-A1.
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                                                             ADE24002 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237
US2003092110-A1.
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Human PRO polynucleotide #237.
US2003203439-A1.
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Human PRO polynucleotide #237.
US2003199062-A1.
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Human PRO polynucleotide #237.
US2003199054-A1.
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23-OCT-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
      (GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 228
ID ADE91215 standard; cl
DE Human PRO polymucleo
PN US200199661-A1.
PD 23-OCT-2003.
PA (GETH ) GENENTECH IN
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Best Local Similarity
RESULT 222
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Best Local Similarity
RESULT 223
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Best Local Similarity
RESULT 226
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Best Local Similarity
RESULT 229
                     Query Match
Best Local Similarity
RESULT 221
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Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207360-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2001199051-A1.
33-OCT-2003.
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Novel human secreted and transmembrane protein PRO1159 CDNA
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Pred. No. 3.3e-75;
                                                                                100.0%; Score 273; DB 12; 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
PRY MATCh 100.0%; SCORE 273; DB 12;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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cDNA encoding human PRO polypeptide #237.
US2003199029-A1.
               #237
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US2003228655-A1.
            cDNA encoding human PRO polypeptide US2003199052-A1. 23-OCT-2003. (GETH ) GENENTECH INC.
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US2003199063-A1.
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US2003207352-A1.
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Human PRO polynucleotide #237.
US2003199060-A1.
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23-OCT-2003.
(GETH ) GENENTECH INC,
100.0%;
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(GETH ) GENENTECH INC.
100.0%;
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ADE95356 standard; cDNA; 713
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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ADG13422 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2003207357-A1.
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06-NOV-2003.
(GETH ) GENENTECH INC.
100.0%;
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(GETH ) GENENTECH INC.
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RESULT 257
ID ADG07855 standard:
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                                                                                        Length 713;
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                                                                                                                                                                                                        Length 713;
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                                                                                                                                                                                                                                           Novel human secreted and transmembrane protein PRO1159 US2003207426-A1.
06-NOV-2003.
(GETH ) GENENTECH INC.
ery Match
st Local Similarity 100.0%; Pred. No. 3.3e-75;
                                                O6-NOV-2003.

(GETH ) GENENTECH INC.

(GETY Match 100.0%; Score 273; DB 12;

lery Match 100.0%; Pred. No. 3.3e-75;
                                                                                                                            ADF98151 standard, cDNA, 713 BP.

Human PRO polynucleotide #237.
US2003207422-A1.
US2003207422-A1.
(GETH) GENENTECH INC.
ery Match
100.0%; Score 273; DB 12;
st Local Similarity 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.

(ery Match

(ery Match 100.0%; Score 273; DB 12;

(ery Match 100.0%; Pred. No. 3.3e-75;
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             ADG20245 standard; cDNA; 713 BP, cDNA encoding human PRO polypeptide #237.US2003207376-A1.
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cDNA encoding human PRO polypeptide #237.
US2003207425-A1.
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Human PRO polynucleotide #237.
US2003207351-A1.
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Human PRO polynucleotide #237.
US2003208055-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
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RESULT 242
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Best Local Similarity
RESULT 240
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Best Local Similarity
RESULT 248
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RESULT 239
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Length 713;
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                                           AUGUB479 standard; CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA
US2203207424-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207389-A1.
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Novel human secreted and transmembrane protein PR01159 cDNA.
US2002207350-A1.
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Novel human secreted and transmembrane protein PRO1159 US2003207427-A1.
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(GETH ) GENENTECH INC.
(ery Match 100.0%; Score 273; DB 12;
ery Match 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
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Arv Match 100.0%; Score 273; DB 12;
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(GETH) GENENTECH INC.

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100.0%; Score 273; DB 12;

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(GETH ) GENENTECH INC.

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Lery Match 100.0%; Pred. No. 3.3e-75;
Score 273; DB 12;
Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
iry Match 100.0%; Score 273; DB 12;
iry Match 100.0%; Pred. No. 3.3e-75;
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Query Match
100.0%; Score 273; DB 12;
Best Local Similarity 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Human PRO polynucleotide #237.
US2003207374-A1.
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Length 713;

Length 713,

Score 273; DB 12; Pred. No. 3.3e-75;

CDNA

Length 713

RESULT 260

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ADG54246 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207416-A1.
06-NOV-2003.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207421-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
                                                                                                     ADG58110 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207363-A1.
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                                                                                                                                                                                                                                        ADG53694 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159
US2003207415-A1.
06-NOV-2003.
(GETH ) GENENTECH INC.
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06-DOV-2003.

(GETH ) GENENTECH INC.

100.0%; Score 273; DB 12;

ATV Match ... 100.0%; Pred. No. 3.3e-75;
         GE-NOV-2003.
(GETH) GENENTECH INC.
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ery Match
ery Match 100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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24-APR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 268
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RESULT 272
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                                                            Best Local Similarity RESULT 267
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US2003207420-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207365-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA,
US2003207428-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA US2003207364-A1.
 Novel human secreted and transmembrane protein PRO1159 cDNA.
                                                                                                                           ADG55350 standard; cDNA; 713 BP. Novel human secreted and transmembrane protein PRO1159 cDNA
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207390-A1.
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US-VCV-2003.

(GETH) GENENTECH INC.

(GETH) GENENTECH INC.

100.0%; Score 273; DB 12;

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100.0%; Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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06-NOV-2003.
(GETH ) GENENTECH INC.
100.0%;
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(GETH ) GENENTECH INC.
"...ch 100.0%;
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US2003207358-A1.
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RESULT 265
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Best Local Similarity
RESULT 263
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Best Local Similarity
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RESULT 259
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ADG82319 standard;
                                                                                                                                                              US2003194778-A1.
                      US2003207356-A1.
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                                                                                                                                                                                 16-OCT-2003
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                                                                         Query Match
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Length 713

Length 713

Length 713

Length 713

Score 273; DB 12; Pred. No. 3.3e-75;

Length 713

Length 713

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Query Match
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                             Length 713;
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                                                                                                                                                                                                                                                                     Length 713;
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                                                                                                                                                                                          ADG56454 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
062003207366-A1.
06-NOV-2003.
                                                                                                                                                                                                                                                                                                            ADH12720 standard, cDNa, 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207378-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207369-A1.
06-NOV-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADG54798 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207367-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADH20155 standard; cDNA; 713 BP.
Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003219856-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                 ADH21115 standard; cDNA; 713 BP.
Human cDNA encoding secreted/transmembrane protein PRO1159
US2003224358-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADG61566 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159
US2003207429-A1.
(GETH ) GENENTECH INC.

3ry Match 100.0%; Score 273; DB 12;

100.0%; Pred. No. 3.3e-75;
                                                                                                                                                 Score 273; DB 12;
Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
(GETY Match 100.0%; Score 273; DB 12;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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                                                                    ADG81215 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003194793-A1.
                                                                                                                                            100.0%;
                                                                                                                                                                                                                                                                   100.0%;
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100.0%;
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100.0%;
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                            ...cr. ,-2003.
...cr. ) GENENTECH I.
..ery Match
Best Local Similarity 1
RESULT 278
ID ADH12720 stand>-
DE Novel humar
PN US2003>-
PD 06-**
                                                                                                                                                                                                                                                    (GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 280

ID ADG6156 Standard; CI
DE Novel human secreted
PN US2003207429-A1.
PD 06-NOV-2003.
PA (GETH ) GENENTECH INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best_Local Similarity
RESULT 282
                          Query Match
Best Local Similarity
RESULT 276
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Best Local Similarity
RESULT 281
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Length 713;
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                                                                                                                                                                            ADG10005 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA. US2004009548-A1.
                                                                                                                                                                                                                                                                                                                     ADII5476 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
06-NOV-2003.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207383-A1.
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NOVEL human secreted and transmembrane protein PRO1159 cDNA.
26-FEB-2004.

(GETH) GENENTECH INC.
100.0%; Score 273; DB 12; Length 713
st Local Similarity 100.0%; pred. No. 3.3e-75;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADG09353 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2004009547-A1.
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                                                                                                                                  Score 273; DB 12;
Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.
iry Match 100.0%; Score 273; DB 12;
iry Match 100.0%; Pred. No. 3.3e-75;
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(GETH ) GENENTECH INC.

(GETH ) GENENTECH INC.

100.0%; Score 273; DB 12;

LETY MATCH IOO.0%; Pred. No. 3.3e-75;
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26-PEB-2004.
26-PEB-2004.
100.0$; Score 273; DB 12;
100.0$; Pred. No. 3.3e-75;
Best Local Similarity 100.0%; Pred. No. 3.3e-75; RESULT 285
                                                                                                                                                                                                                                                                                 Score 273; DB 12;
Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
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Pred. No. 3.3e-75;
                                       ADI81262 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237 US2003207361-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  cDNA encoding human PRO polypeptide #237
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06-NOV-2003,
(GETH ) GENENTECH INC.
100.0%; S.
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                                                                                                                                100.0%;
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15-JWN-2004.
(GETH ) GENENTECH INC.
Match 'Totter 100.0%;
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15-JAN-2004.
(GETH ) GENENTECH INC.
"" Match '''rity 100.0%;
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                                                                                           06-NOV-2003,
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            06-NOV-2003.
(GETH ) GENENTECH INC.
                                                                                                                                Query Match
Best Local Similarity
RESULT 286
                                                                                                                                                                                                                                                                                        Best_Local Similarity RESULT 287
                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 288
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 289
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RESULT 292
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Score 75.2; DB 2;
Pred. No. 3.1e-13;
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02-JUL-1998.
(TEXA ) UNIV TEXAS SYSTEM.
STY Match
Et Match
St Local Similarity 77.2%; Pred. No. 3.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABZ24043 standard; DNA; 2298 BP.
Mouse Dectin-1 polypeptide encoding DNA
Molose6945-A2.
05-DBC-2002.
(ISIS-) ISIS INNOVATION LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABZ24041 standard; DNA; 1606 BP.
Human Dectin-1 polypeptide encoding DNA
WO200296945-A2.
                                                                                                                         ABQ66733 standard, cDNA, 1281 BP.
Human polynucleotide SEQ ID NO 223.
US2002042386-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      CDNA; 2298 BP
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   28.4%;
73.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                02-JUL-1998.
(TEXA ) UNIV TEXAS SYSTEM.
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Best Local Similarity
                                                                                                                                                                                  11-APR-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Mouse dectin-1 cDNA.
WO9828332-A2,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
                                                                                   Local Similarity
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Best Local Similarity
RESULT 304
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ID AAV42548 standard,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAV42551 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
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                                                                     Query Match
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RESULT 305
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36094 WO200157182-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADF60937 standard; cDNA; 1018 BP.
Human long Schering dendritic cell membrane protein 4 (SDCMP4) cDNA
US2003162955-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        03-OCT-ZUUG.
(NOVS) NOVARTIS AG.
(NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH.
51.3%; Score 140; DB 10; Length 1153;
2TY Match 51.3%; Pred. No. 2e-33;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   51.3%; Score 140; DB 10; Length 1018; 100.0%; Pred. No. 1.9e-33;
                                                                                                         Length 713;
                                                                                                                                                                                                                                                               100.0%; Score 273; DB 12; Length 713; 100.0%; Pred. No. 3.3e-75;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 100.0%; Score 273; DB 4; Length 5709;
ery Match 100.0%; Pred. No. 7e-75;
                                                                                                                                                                                                                                                                                                                                                                                                                    Length 713;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 741;
                                                               UL-MAR-2004.
(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
ery Match
ery Match 100.0*; Pred. No. 3.3e-75;
                                                                                                                                                                                                                                                                                                                                                                     USZUCZ-VOR.
22-APR-2004.
(GETH) GENENTECH INC.
ery Match 100.0%; Score 273; DB 12;
ery Match 100.0%; Pred. No. 3.3e-75;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 140; DB 2; ]
Pred. No. 1.7e-33;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               51.3%; Score 140; DB 2; 3100.0%; Pred. No. 1.9e-33;
             ADM27837 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2004048333-A1.
                                                                                                                                                              ADM42561 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2004058424-A1.
                                                                                                                                                                                                                                                                                                                     ADM28423 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237.
US2004077064-A1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAZ07533 standard; cDNA; 1018 BP.
Human SDCMP4 polypeptide encoding cDNA.
W09947673-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABV73351 standard; cDNA; 1153 BP.
LLR-2734-2 polypeptide encoding cDNA.
WO200277216-A2.
03-OCT-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAX01260 standard; cDNA; 741 BP.
Human DC3' protein coding sequence.
JP11001497-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAK81282 standard; DNA; 5709 BP
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(FAKE) TAKEDA CHEM IND LTD.

(Ery Match ....1arity 100.0%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           23-SEP-1999.
(SCHE) SCHERING CORP.
Query Match 5
                                                                                                                                                                                                                          25-MAR-2004.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (CHAL/) CHALUS L.
(QUAN/) QUAN A B.
(BATE/) BATES E B M.
(GORM/) GORMAN D M.
(SAEL/) SAELAND S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 301
D ABV7331 standard; cD
DE LLR-U24-2 polypeptide
PN WO200277216-A2.
PD 03-0CT-2002.
PA (NOVS ) NOVARTIS-ERFII
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 302
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 300
                                                                                                           Query Match
Best Local Similarity
RESULT 295
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 298
                                                                                                                                                                                                                                                                                   Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 297
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                28-AUG-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
RESULT 294
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Human cDNA encoding a novel extracellular matrix protein, Seq 1D No 223. WO200155368-Al. 02-AUG-2001. (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAK81284 standard; DNA; 336 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36096.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 2298;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 1281;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 2298
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 528;
                                                                                                                                                                                                                                                                                                           Length 1281;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 1606;
                                                                                         Length 1281;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 138;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Ouery Match 41.4%; Score 113; DB 4; Length 336; Best Local Similarity 100.0%; Pred. No. 3.7e-25;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WOLDEC-2002.

0S-DEC-2002.

(ISLS-) ISIS INNOVATION LTD.

51.3%; Score 140; DB 8; I

(ery Match 100.0%; Pred. No. 2.2e-33;
                                                                                                                                                                                                                                                                                                                                                                                                                                USACUSTONS.
27-MAR-2003.
(HUMA-) HUMAN GENOME SCI INC.
61.3%; Score 140; DB 10;
ery Match 51.3%; Score 140; DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABY73363 standard; CDNA; 138 BP.

LLR-0724-stalk peptide encoding cDNA.

WO200277216-A2.

O3-OCT-2002.

(NOVS) NOVARTIS AG.

(NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH.

Query watch 50.5%; Score 138; DB 10;

Best Local Similarity 100.0%; Pred. No. 3.9e-33;
                                                                   SCI INC.
51.3%; Score 140; DB 4; 1
100.0%; Pred. No. 2.1e-33;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 77.6; DB 2;
Pred, No. 9.4e-14;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 28.4%; Score 77.6; DB B; 73.1%; Pred. No. 9.4e-14;
                                                                                                                                                                                                                                                                                                             51.3%; Score 140; DB 6; 100.0%; Pred. No. 2.1e-33;
                                                                                                                                                                                                                                                                                                                                                                                                #5
                                                                                                                                                                                                                                                                                                                                                          Human cDNA from extracellular matrix gene 66
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Length 2086;

Length 2086;

PN

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ABK91073 standard; DNA; 5144 BP.
Gene encoding Arabidopsis thaliana SOS2 serine/threonine protein kinase.
VS2002095032-Al.
                                                                                                                                                                                                                                                                                      Human musculoskeletal system-associated genomic DNA - SEQ ID 3431.
US2004009488-A1.
                                                               ABX60054 standard; cDNA; 2086 BP. cDNA encoding novel human musculoskeletal system antigen #2398. US2002147140-A1.
                                                                                                                                                                                                                                                                                                                                                                  DB 12; Length 2086;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AG.
12.3%; Score 33.6; DB 6; Length 9646;
48.9%; Pred. No. 9.9;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 6; Length 5144;
        12.6%; Score 34.4; DB 4; 57.4%; Pred. No. 3.2;
                                                                                                                                                                                                          DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                    D ABL99813 standard; cDNA; 4045 BP.

E Human secretory polynucleotide (sptm) 68.

N W0200220756-A2.

I 14-MAR-2002.

A (INCY-) INCYTE GENOMICS INC.

Query Match

Best Local Similarity 57.4%; Pred. No. 4.1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           12.5%; Score 34; DB 6; 52.9%; Pred. No. 20;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    GGNO-) GENOME THERAPEUTICS CORP.

(GENO-) GENOME THERAPEUTICS CORP.

12.5%; Score 34.2; DE

12.5%; Pred. No. 2.5;
                                                                                                                                                                                                        12.6%; Score 34.4; Di 57.4%; Pred. No. 3.2;
                                                                                                                                                                                                                                                                                                                                        (HUMA-) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 57.4%; Pred. No. 3.2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match 12.4%; Score 33.8; DE
Best Local Similarity 50.3%; Pred. No. 6.8;
RESULT 328
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             12.3%; Score 33.6; DI 56.2%; Pred. No. 2.9;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              12.5%; Score 34.2; 1
50.3%; Pred. No. 15;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          RESULT 324

ID ADC92020 standard; DNA; 696 BP.

DE E. faccium DNA sequence SEQ ID 1647.

PN US6583275-81.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAS98600 standard; DNA; 140036 BP.
Human genomic DNA for PHIP/NDRP.
WQ200185785-A2.
15-NOV-2001.
(ROZA), ROZAKIS-ADCOCK M.
(FARH/) FARHANO-FALLAH J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABN77024 standard; cDNA; 321 BP.
Human ORF1971 cDNA, SEQ ID NO:3941.
WO200190366-A2.
                                                                                                                                                                                                                                                                 ADJ30804 standard; DNA; 2086 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DNA; 9646
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             18-JUL-2002.
(UYAR-) UNIV ARIZONA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      29-NOV-2001.
(CURA-) CURAGEN CORP.
                                                                                                                         10-OCT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                03-JAN-2002.
(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
        Query Match
Best Local Similarity
RESULT 321
                                                                                                                                                                                                                    Best Local Similarity
RESULT 322
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 326
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 325
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 329
ID ABL33689 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
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                                                                                                                                                                                                                                                                                                                           15-JAN-2004
                                                                                                                                                                                                          Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AALJ7066 standard; DNA; 2086 BP.
Human musculoskeletal system related polymucleotide SEQ ID NO 3431.
WC200155367-A1.
                                                                                                                                                                                                                                                                                                                     Human musculoskeletal system related polynucleotide SEQ ID NO 696 WO200155367-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human musculoskeletal system-associated contig DNA - SEQ ID 696 US2004009488-A1.
                                                                                                                                                                                   Score 37.8; DB 6; Length 110000;
Pred. No. 1.2;
                                                                                                                                                                                                                                           Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABX58342 standard; cDNA; 930 BP.
cDNA encoding novel human musculoskeletal system antigen #686.
US202147140-A1.
10-0CT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     12.6%; Score 34.4; DB 12; Length 1446; 57.4%; Pred. No. 2.8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             15-JAN-2004.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match
ery Match
(ery Match 57.4%; Pred. No. 2.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 1969;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 1446;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 12.6%; Score 34.4; DB 8; Length 930; 57.4%; Pred. No. 2.4;
                                                                                                                                                                                                                                                                                                                                                                                                     Length 930
                                                                                                                           Length 85
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AEK43739 standard; cDNa; 1446 BP.
DNA encoding novel central nervous system protein #319.
WO200155318-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  07-MAY-2003.

(HELI-) HELIX RES INST.

(REAS-) RES ASSOC BIOTECHNOLOGY.

Query Match

12.6%; Score 34.4; DB 10;

Best Local Similarity 57.4%; Pred. No. 3.1;
                  AAC24137 standard; cDNA; 85 BP.
Human secreted protein 5' EST, SEQ ID NO: 28212.
EP1033401-A2.
06-SEP-2000.
(GEST ) GENSET.
                                                                                                                                                                                                                                           Score 37.8; DB 6;
Pred. No. 1.2;
                                                                                                                                                                                                                                                                                                                                                MOZOCE-2001.

02-ADG-2001.

(HUMA-) HUMAN GENOME SCI INC.

12.6%; Score 34.4; DB 4;

lery Match

12.6%; Score 34.4; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     12.6%; Score 34.4; DB 4; 57.4%; Pred. No. 2.8;
                                                                                                                       Score 74; DB 3;
Pred. No. 3.9e-13;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADID4126 standard; cDNA; 1446 BP. cDNA encoding novel human protein seq id 329. US2004018969-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADB63593 standard; cDNA; 1969 BP.
Human cDNA encoding clone THYMU20034790.
EP1308459-A2.
                                                                                                                                                                                                                                                                                                    ВP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADJ28069 standard; DNA; 930 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                     27.18;
97.48;
                                                                                                                                                                                                                                           13.8%;
53.8%;
                                                                                                                                                                                                                                                                                                    CDNA; 930
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ID AAL37066 standard; DNA; 2086
DE Human musculoskeletal system
WO200155367-A1.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                 13.8%;
                                                                                                                                                                                                    53.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PD 29-JAN-2004.
PA (ROSE/) ROSEN C A.
PA (RUBE/) RUBEN S M.
PA (BARA/) BARASH S C.
Query Match
Best Local Similarity
RESULT 319
                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
RESULT 315
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 318
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 316
                                                                                                                     Query Match
Best Local Similarity
RESULT 312
                                                                                                                                                                               Query Match
Best Local Similarity
RESULT 313
                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 314
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
                                                                                                                                                                                                                                                                                                AAL35354 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             RESULT 317

ID ABK4373

DE DNA enc

PN WO20015

PD 02-AUG-

PA (HUMA-)
RESULT 311
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Length 110000;

Length 140036;

Length 321;

PA PE

Length 696;

Length 4045;

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AAK74053 standard; DNA; 39567 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:28865.
WO200157182-A2.
                                                                                                                                                                                                                                               ABZ74429 standard; DNA; 39567 BP.
Secreted protein gene 286 genomic fragment HRGBD54, SEQ ID NO:1576.
WO200277013-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DB 10; Length 39567;
                                                                                                                                                                                                                                                                                                                                                  Length 39567;
                                                                                                                                                                                        Length 39567;
(ELIT-) ELITRA PHARM INC.

ery Match 11.9%; Score 32.6; DB 8; Length 1209;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   22-MAY-2003.

4 (INCY-) INCYTE GENOMICS INC.

Query Match

Best Local Similarity 51.4%; Pred. No. 16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         11.7%; Score 32; DB 11; Length 1414; 51.4%; Pred. No. 16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 1432;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11.7%; Score 32; DB 10; Length 695; 48.6%; Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                                                       ABZ67986 standard; DNA; 39567 BP.
Human secreted protein encoding genomic DNA SEQ ID NO 1509
WO200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 366;
                                                                                                                                                                                        DB 4;
                                                                                                                                                                                                                                                                                                                                                  DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     11;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human colon cancer cell expressed cDNA #1288.
US2002155438-A1.
24-OCT-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. ery Match

11.7%; Score 32; DB 5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB
16;
                                                                                                                                         (HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

11.9%; Score 32.4; D

ery Match

11.9%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity 48.6%; Pred. No. 12; RESULT 344
ID ADM23365 standard; DNA; 1414 BP.
DE Human novel protein NOV22b coding sequence. PN WO2003064628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human novel protein NOV22a coding sequence. WO2003064628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABV06453 standard; cDNA; 366 BP.
Human prostate expression marker cDNA 6444.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                11.9%; Score 32.4; I
50.0%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (HUMA-) HUMAN GENOME SCI INC.
ry Match 11.9%; Score 32.4; I
Local Similarity 50.0%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 32;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABL58125 standard; CDNA; 1889 BP.
Human RNA helicase 12 coding sequence.
CN1331331-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DNA; 1529 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CDNA; 695 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.7%;
51.4%;
                                                                                                                                                                                                                                                                                                                               (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADC14278 standard; DNA; 15:
Human enzyme ENZM-31 gene.
WO2003042357-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (SIMP/) SIMPSON A J G. (NETO/) NETO E D. (BREN/) BRENTANI R R.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    07-AUG-2003.
(CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          07-AUG-2003.
(CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 345
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best_Local Similarity
RESULT 346
                                                                                                                                                                                                                                                                                                                                                                Local Similarity
                                Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ACD92876 standard;
                                                                                                                                                                                                                                                                                                           03-OCT-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      03-OCT-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
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                                                                                                                                                                                          Ouery Match
                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                        RESULT 341
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADO47190 standard, DNA, 200000 BP.

DNA sequence of a human immunoglobulin heavy chain variable region.
WO2004029249-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 200000
                                                                                                                                 Length 18218;
                                                                                                                                                                                                                                                                                                             Length 1990;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Human reproductive system related antigen DNA SEQ ID NO: 8316. WO200155320-A2. 02-AUG-2001. (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (BPIG-) EPIGENOMICS AG. 12.0%; Score 32.8; DB 6; Length 9832; Pred. No. 18;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           12.0%; Score 32.8; DB 6; Length 435; 57.4%; Pred. No. 5.8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   12.1%; Score 33; DB 4; Length 32176; 46.1%; Pred. No. 24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 9; Length 489,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         12.1%; Score 33; DB 3; Length 2928; 51.0%; Pred. No. 9.9;
                                                   Human immune system associated gene SEQ ID NO: 1921.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAC39967 standard; DNA; 2928 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 26555
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABN16359 standard; cDNA; 435 BP.
Human ORFX polynucleotide sequence SEQ ID NO:1195.
WO200192523-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                         GB-APR-2004.
(CENT-) CENTENARY INST CANCER MEDICINE & CELL BI.
(CENT-) CENTENARY INST CANCER MEDICINE & CELL BI.
ery Match
12.2%; Score 33.4; DB 12;
cr Local Similarity 55.7%; Pred. No. 34;
                                                                                                                                                                                                                                                                                                         12.2%; Score 33.4; DB 10; 54.5%; Pred. No. 6.5;
                                                                                                                                 DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL32657 standard; DNA; 9832 BP.
Human immune system associated gene SEQ ID NO:
WO200200928-A2.
                                                                                                            AG.
12.3%; Score 33.6; 1
50.6%; Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            11.9%; Score 32.6; 48.9%; Pred. No. 7;
                                                                                                                                                                                                            Human cDNA encoding clone PLACE60175640.
BP1308459-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ACA28755 standard; DNA; 1209 BP.
Prokaryotic essential gene #10412.
WO200277183-A2.
03-OCT-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DNA; 32176 BP.
                RESULT 330
ID ABL33948 standard; DNA; 18218 BP.
                                                                                                                                                                                          ADB62949 standard; cDNA; 1990 BP
                                                                                                                                                                                                                                                                      (REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  .T 337
ACH22923 standard; cDNA; 489 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human adult ovary cDNA #1303
US2003073623-A1.
17-APR-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (DRMA/) DRMANAC R T.
(LABA/) LABAT I.
(STAC/) STACHE-CRAIN B.
(DICK/) DICKSON M C.
(JONE/) JONES L W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    06-DEC-2001.
(CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
RESULT 337
RESULT 337
DE Human adult covary CDN
N US2003073623-A1.
PA (DRMA) DRMANAC R T.
PA (STACA) STACHE-CRAIN
PA (STACA) STACHE-CRAIN
PA (STACA) STACHE-CRAIN
PA (JONE/) JONES L W.
                                                                                        03-JAN-2002.
(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 333
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity
RESULT 334
ID AAL05628 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Ouery Match
Best Local Similarity
RESULT 336
                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 332
                                                                                                                                                   Best Local Similarity
RESULT 331
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 335
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   EP1033405-A2.
                                                                                                                                                                                                                                                 07-MAY-2003
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Query Match

Query Match

Query Match

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AAZS6348 standard; DNA; 1665 BP.
Escherichia coli flagellin protein nucleotide sequence SEQ ID NO:18.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11.6%; Score 31.6; DB 12; Length 161671;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 6; Length 14307;
                                                                                                                                                                                                                                                                 ABX04178 standard, cDNA, 2680 BP.
Human mRNA differentially expressed in mesenchymal cells
WO200271927-A2.
                                                                                     DB 10;
                                                                                                                                                                                                                     DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                  AAQ81792 standard, DNA; 8478 BP.
B. subtilis biotin operon and flanking sequences.
EP635572-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 3;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human metastasis associated gene SEQ ID NO: 180 WO200177376-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       LT 364
ABL32728 standard, DNA; 14307 BP.
ABL32728 standard, DNA; 14307 BP.
Human immune system associated gene SEQ ID NO:
W0200200928-A2.
03-0028-A2.
(EPIG.) EPIGENOMICS AG.
(EPIG.) EPIGENOMICS AG.
11.6%; Score 31.6; DB 6
                                                                                                                                                                                                                                                                                                                                                      DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB
                                                                                                                                                                                                                                                                                                                                                                                                                                             (HOFF ) HOFFMANN LA ROCHE & CO AG F. ery Match 11.6%; Score 31.8; I for a familarity 56.1%; Pred. No. 35;
                                                                                    Score 31.8; I
Pred. No. 16;
                                                                                                                                                                                                                     11.6%; Score 31.8; 1 56.1%; Pred. No. 23;
                                                                                                                                                                                                                                                                                                                                                    11.6%; Score 31.8; I 56.1%; Pred. No. 23;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    11.6%; Score 31.6; 150.7%; Pred. No. 12;
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Pred. No. 22;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 31.6; I
Pred. No. 34;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 31.8;
Pred. No. 88;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAV/5482 standard; DNA; 299 BP.
Staphylococcus aureus contig SEQ ID #1171
EP786519-11
                                                                                                                                                                                                                                                                                                                 19-SEP-2002.
(BGHM ) BRIGHAM & WOMENS HOSPITAL INC.
   DEUT KREBSFORSCHUNGSZENTRUM.
UNIV LUDWIG MAXIMILIANS.
HAFERLACH T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DNA; 161671 BP
                                                                                                                   RESULT 357

ID ACA92148 standard; DNA; 2648 BP.
DE DNA encoding human PMMM-33.
PN WO2003031939-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL34627 standard; DNA; 5539 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  30-JUL-1997.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                     11.6%;
58.7%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     11.6%;
54.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.6%;
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                                                                                                                                                                                  17-APR-2003.
(INCY-) INCYTE GENOMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              0 03-JUL-2003.
(VITI-) VITIVITY INC.
Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WO9961458-A1.
02-DEC-1999.
(UNSY ) UNIV SYDNEY.
                                                                                                                                                                                                                               Best Local Similarity RESULT 358
                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
RESULT 359
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADH30232 standard;
Human PLOD2 DNA.
US2003124535-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
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                                                                                                   Best Local Similarity
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Best Local Similarity
                                                     (SCHO/) SCHOCH C. (KERN/) KERN W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    EP786519-A2.
                                                                                                                                                                                                                     Ouery Match
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                                                                                       Query Match
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                    (UYLU-)
(HAFE/)
      (DEKR-)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 361
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     RESULT 364
                                                                                                                                                                                                                                                                                                  Length 2114;
                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 2183;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 713;
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                                     Length 1889;
                                                                                                                                                                   Length 1931;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 2504;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 2265;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                AMSW/446 standard; cDNA; 2265 BP.
DNA encoding novel human diagnostic protein #23250
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 tumour protein, SEQ ID No 1016
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AEK55395 standard; cDNA; 713 BP.
Human colon cancer-associated cDNA, SEQ ID No 865.
WO200212280-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  . 9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.6%; Score 31.8; DB 6; 56.1%; Pred. No. 14;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAH33427 standard; cDNA; 851 BP.
Human colon cancer antigen encoding cDNA SEQ ID
WC200122920-A2.
                                                                                                                                                                 Score 32; DB 4;
Pred. No. 18;
                                                                                                                                                                                                                                                                                                    Score 32; DB 12;
Pred. No. 18;
                                                                                                                                                                                                                                                                                                                                                                                                                                 11.7%; Score 32; DB 11; 51.4%; Pred. No. 18;
                                  11.7%; Score 32; DB 6; 47.1%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  7 alpha hydroxylase sequence encoding sequence WO200231111-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.7%; Score 32; DB 6;
51.4%; Pred. No. 19;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    田
                                                                               AAD09939 standard; cDNA; 1931 BP.
Human drug metabolising enzyme (DME-4) cDNA.
W0200151638-A2.
19-JUL-2001.
(INCY-) INCYTE GENOMICS INC.
ETY Match
Struct 17%; Score 32; DB 4
st Local Similarity 51.4%; Pred. No. 18;
                                                                                                                                                                                                                                                                                                                                                                Human novel protein NOV22c coding sequence WO2003064628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.6%; Score 31.8; 56.1%; Pred. No. 13;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 32;
Pred. No.
16-JAN-2002.
(HODE-) BODE GENE DEV CO LTD SHANGHAL.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADF82256 standard; DNA; 983 BP.
Leukaemia-related DNA sequence #2812.
WO2003039443-A2.
15-MAY-2003.
                                                                                                                                                                                                               ADNO4961 standard; cDNA; 2114 BP. Antipsoriatic cDNA sequence #695. WO2004028479-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  CDNA; 2504 BP
                                                                                                                                                                                                                                                                                                                                               ADM29367 standard; DNA; 2183 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                CDNA; 506 BP.
                                                                                                                                                                                                                                             WO2004020...
08-APR-2004.
(GETH ) GENENTECH INC.
"artch 11.7%; Sc
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(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.7%;
51.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 51.4%;
                                                                                                                                                                                                                                                                                                                                                                       J.2003.
J.2003.
J.RA-) CURAGEN CON.
J.GY MAtch
Best Local Similarity
RESULT 351
ID AAS87446 stand*
DE DNA encodir
PN W020017*
PD 11.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         "...A-) HUMAN GENOM.
"...a-) HUMAN GENOM.
Best Local Similarity RESULT 356
ID ADF82256 star.
DE Leukaemi-
PN WO?"
                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 350
ID ADM29367 standard; DN
DE Human novel protein N
PN WO2003064658-A2.
PD 07-AUG-2003.
PA (CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                               07-AUG-2003.
(CURA-) CURAGEN CORP.
                                              Best Local Similarity RESULT 348
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABK45465 standard; c cDNA encoding colon WO200212328-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 355
                                                                                                                                                                 Query Match
Best Local Similarity
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Best Local Similarity
RESULT 352
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABQ60988 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               14-FEB-2002
                                  Query Match
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                                                                                                                                                                                                RESULT 349
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Length 110000;

Length 8478;

Length 5539;

9

PA PA

Length 1665

Length 299

Length 2648;

PN PN

Length 2680;

9

Length 983;

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DB 11; Length 3142;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 12; Length 3142;
                                                                                                                                             DB 12; Length 1811;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 2; Length 3142;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADP13425 standard; DNA; 3142 BP. Renal carcinoma differentially expressed gene #161 WO2004048933-A2.
                                                                                                                                                                                                 AAS41019 standard; cDNA; 1854 BP. cDNA encoding novel human enzyme polypeptide #235.
WO200155301-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match 11.4%; Score 31.2; DB 2;
Best Local Similarity 60.7%; Pred. No. 36;
                                                                                                                                                                                                                                                                                                 DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human BEC/LEC-related gene sequence SeqID470.
WO2003080640-A1.
             ADI53814 standard; cDNA; 1811 BP. cDNA encoding novel human protein seq id 17. US2004018969-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity 66.2%; Pred. No. 37; RESULT 380
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (INCY-) INCYTE GENOMICS INC.
ry Match 11.4%; Score 31.2; I
t Local Similarity 66.2%; Pred. No. 36;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 31.2; Pred. No. 37;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 31.2; I
Pred. No. 37;
                                                                                                                                                                                                                                                                                               11.4%; Score 31.2; 1 66.2%; Pred. No. 31;
                                                                                                                                             Score 31.2; |
Pred. No. 31;
                                                                                                                                                                                                                                                                                                                                                                                                                             (WHED ) WHITEHEAD INST BIOMEDICAL RES. (MILL-) MILLENNIUM PHARM INC.
                                                                                                                                                                                                                                                                                                                                                   AAZ00354 standard; DNA; 2885 BP.
Nucleotide sequence of human hsFATP6.
WO9936537-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADD69636 standard; cDNA; 2899 BP.
Human REMAP cDNA - SEQ ID 65.
WO2003048305-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADI31764 standard; cDNA; 3142 BP.
Human cDNA #1090.
US6607879-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 379
ID AAQ82847 standard; DNA; 3142 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DNA; 3199 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             0 02-OCT-2003.
1 (LUDW-) LUDWIG INST CANCER RES.
A (LICK) ) LICENTIA LTD.
11.4%; Sc
                                                                                                                                                                                                                                                            02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
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66.2%;
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                                                                                                                                               11.4%;
66.2%;
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(OKLA ) UNIV OKLAHOMA STATE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (AMHP) WYETH.
(TWIN) TWINE N C.
(TREP) BURCZYNSKI M E.
(TREP) TREPICCHIO W L.
(DORN) DORNER A.
(STOV) STOVER J A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               19-AUG-2003.
(INCY-) INCYTE CORP.
                                                                     29-JAN-2004.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADN95547 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
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Best Local Similarity
RESULT 376
                                                                                                                                                                                                                                                                                                                Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           P-selectin gene WO9506118-Al.
                                                                                                                                                                                                                                                                                                                                                                                                               22-JUL-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              12-JUN-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
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RESULT 382
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 378
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABX63053 standard; cDNA, 1072 BP.

Human cDNA #53 differentially expressed in activated vascular tissue.
US2002137081-A1.

(BAND/) BANDMAN O.
                                                                                                                                                                                                       NO 59
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAK58497 standard; cDNA; 559 BP.
Human immune/haematopoietic antigen encoding cDNA SEQ ID NO:3557.
W0200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ACA04691 standard; CDNA; 1760 BP.

CDNA encoding human membrane associated protein fragment #139.
US6492505-81.
10-DEC-2002.
(INCY-) INCYTE GENOMICS INC.

ETY MATCH MATCH 11.4*; Score 31.2; DB 8; Length 1760; st Local Similarity 66.2*; Pred. No. 30;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 4; Length 1811;
                                                                                                                                Length 6156;
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                                                                                                                                                                                                                                                                                 Length 6156
                                                                                                                                                                  ABL49359 standard; DNA; 6156 BP.
Human polynucleotide associated with DNA replication SEQ ID :
W0200177377-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 536;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 4; Length 559
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABL45638 standard; DNA; 1202 BP.
Human ubiquitin protein ligase B3A gene fragment SEQ ID NO:
W020192582-A1.
(GENA-) GENAISSANCE PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 536
                                                                                                                                                                                                                                                                                                                                                       Human brain expressed single exon probe SEQ ID NO: 10537 WO200157275-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADN#3427 standard; cDNA; 1811 BP.
DNA encoding novel central nervous system protein
NO200155318-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 8;
                                                                                                                                11.5%; Score 31.4; DB 6; 61.7%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 6;
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   54.2%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABS36095 standard; DNA; 536 BP.
Human liver single exon probe, SEQ ID No 11085.
WO200157273-A2.
                                ABL92288 standard; DNA; 6156 BP.
Chemically treated DNA repair gene fragment#49
WO200181622-A2.
(EPIG-10NOY-2001.
                                                                                                                                                                                                                                                            AG.
11.5%; Score 31.4; DB
61.7%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WACALON OB-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
(ELY Match

Score 31.2; E
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                                                                                                                                                                                                                                                                                                                                                                                    WOLD OF AUG 2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MALCh
11.4%; Score 31.2; I
ery Match
50.7%; Pred. No. 20;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
(SETY MALCh 11.4%; Score 31.2; I
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.4%; Score 31.2;
51.4%; Pred. No. 26;
                                                                                                                                                                                                                                                                                                                                         AAK10546 standard; DNA; 536 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
RESULT 370
ID AAK58497 standard; CL
DE Human immune/haematop
PN WO200157182-A2.
PD 09-AUG-2001.
PA (HUMA-) HUMAN GENOME
                                                                                                                                                                                                                                              18-OCT-2001.
(EPIG-) EPIGENOMICS
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Best Local Similarity
RESULT 373
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Best Local Similarity
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Best Local Similarity
RESULT 369
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Best Local Similarity
RESULT 372
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Best Local Similarity RESULT 366
                                                                                                                                                 Best Local Similarity
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Query Match

ID DE PN

Length 2899;

Length 2885;

Length 1854;

Query Match

Query Match

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(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 394
                                                                        Best Local Similarity RESULT 393
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
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(HYSE-) HYSEQ INC.
         WO200170979-A2.
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                                                             Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAK73896 standard; DNA; 396 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:28708
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL07258 standard; cDNA; 19754 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 16256.
W0200171042-A2.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ID ABN83124 standard; DNA; 172637 BP.
DE Human voltage-activated ion channel transporter protein gene.
Query Match 11.44; Score 31.2; DB 6; Length 172637;
Best Local Similarity 54.38; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                   ABL28550 standard; DNA; 4324 BP. Drosophila melanogaster genomic polymucleotide SEQ ID NO 37123 W0200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 19754;
                                                                                                                                 Length 3246;
                                                                                                                                                                                                                                                                                                                                                                                                          11.4%; Score 31.2; DB 4; Length 4324; 58.7%; Pred. No. 42;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 6074;
                                                                                                                                                                                                                                                                     Length 3660;
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(HUMA-) HUMAN GENOME SCI INC.
(ery Match 11.4%; Score 31; DB 4; Length 396; ery Match 50.3%; Pred. No. 20;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  77-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL33065 standard; DNA; 6074 BP.
Human immune system associated gene SEQ ID NO: 1038
WO200200928-A2.
                                                                                   14-201.
14-201.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
11.4%; Score 31.2; DB 4;
or !ocal Similarity 66.2%; Pred. No. 38;
                                                                                                                                                                                                                                                                     DB 9;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11.4%; Score 31.2; DB 6; 52.3%; Pred. No. 47;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         11.4%; Score 31.2; DB 4; 58.7%; Pred. No. 73;
                                                        Human cervical cancer marker nucleic acid 3857. WO200142467-A2.
                                                                                                                                                                   AAD57202 standard, DNA, 3660 BP.
Rice homologue of petunia restorer (Rhpr2) DNA, WQ2003057859-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11.4%; Score 31; DB 5; 52.8%; Pred. No. 21;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                11.4%; Score 31.2; 53.8%; Pred. No. 40;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            11.4%; Score 31; DB 52.8%; Pred. No. 20;
   66.2%; Pred. No. 38;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADL62560 standard; DNA; 397 BP.
Human ovarian cancer DNA marker #20772.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADI76991 standard; DNA; 400 BP. Human ovarian cancer DNA marker #9733.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADI70673 standard; DNA; 400 BP.
Human ovarian cancer DNA marker #3415.
WO200170979-A2.
                                        AAH72583 standard; cDNA; 3246 BP
                                                                                                                                                                                                                             17-JUL-2003.
(CORR ) CORNELL RES FOUND INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                             Query Match
Best Local Similarity
RESULT 384
Best Local Similarity
RESULT 383
                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 385
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RESULT 388
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
RESULT 392
                                                                                                                                                                                                                                                                                                                                                                                          (PEKE ) PE CORP NY.
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Best Local Similarity
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ABZ09989 standard; DNA; 4233 BP.
Haematopoietic cell proliferation disorder related DNA sequence #129.
WO200277272-A2.
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Haematopoietic cell proliferation disorder related DNA sequence #275.
WO200277272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABZ10093 standard, DNA; 4233 BP.
Haematopoietic cell proliferation disorder related DNA sequence #233 WO200277272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABZ10239 standard; DNA; 4233 BP.
Haematopoietic cell proliferation disorder related DNA sequence #379.
WO200277272-A2.
                                                                                                                                                                                                                                            AAK59117 standard; cDNA; 519 BP.
Human immune/haematopoietic antigen encoding cDNA SEQ ID NO:4177.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 1885;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAS61454 standard; DNA; 4233 BP.
Human gene regulation-associated gene oligonucleotide #409.
WO200177375-A2.
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                                          Length 400;
                                                                                                                                                                                          Length 494;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 823;
                                                                                                                                                                                                                                                                                                                                         Length
                                                                                                                                                                                                                                                                                                                                                                                           AAS92837 standard; cDNA; 823 BP.
DNA encoding novel human diagnostic protein #28641.
WO200175067-A2.
                                                                                              ADL42227 standard; DNA; 494 BP.

Human ovarian cancer DNA marker #16117.
27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
EY MARCH 11.4%; Score 31; DB 5; ELocal Similarity 52.8%; Pred. No. 22;
77-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery March 11.4%; Score 31; DB 5;
er Incal Similarity 52.8%; Pred. No. 21;
                                                                                                                                                                                                                                                                                          11.4%; Score 31; DB 3; 52.8%; Pred. No. 36;
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8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       11.4%; Score 31; DB 5; 48.6%; Pred. No. 27;
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9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ..
8
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48;
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48;
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48;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human secreted protein cDNA sequence #20.
WO200055201-A1.
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Pred. No.
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Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 31;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAC59176 standard; cDNA; 1885 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 21-SEP-2000.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Wozorz...
18-001-2001.
(EPIG-) EPIGENOMICS AG.
11.4%;
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49.1%;
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(EPIG-) EPIGENOMICS AG.
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Query Match

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SEQ ID 3487
                                                                                                                                                                                                                                  ABX60110 standard; CDNA; 32248 BP. CDNA encoding novel human musculoskeletal system antigen #2454. US2002447140-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         11.4%; Score 31; DB 8; Length 139308; 51.0%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ) 20-NOV-2003.

4 (PHAS-) PHASE-1 MOLECULAR TOXICOLOGY INC.

Query Match 11.3%; Score 30.8; DB 12; Length 705;

Best Local Similarity 50.0%; Pred. No. 29;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           15.JAN-2004.

(HUMA-) HUMAN GENOME SCI INC.

(ery Match 11.4%; Score 31; DB 12; Length 32248;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Score 30.8; DB 5; Length 450;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  705;
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                                                                                                                                                                                                Length 32248;
                       Length 32248;
                                                                                                                                                                                                                                                                                                                                                                           Length 32248;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AUG45603 standard; DNA; 705 BP.
Liver inflammatory predictive gene related DNA sequence.
W0200395624-A2.
                                                                                                                                                                                                                                                                                                                                                                                                           мыстивей standard; DNA; 32248 BP.
Human musculoskeletal system-associated genomic DNA
US2004009488-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5;
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                                                                                                                                                                                                                                                                                                                                                                           Score 31; DB 8;
Pred. No. 1e+02;
                       11.4%; Score 31; DB 5; 50.3%; Pred. No. 1e+02;
                                                                                                                                                                                                Score 31; DB 8;
Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

11.3%; Score 30.8; DB

ery_match 48.3%; Pred. No. 23;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (PHAS-) PHASE-1 MOLECULAR TOXICOLOGY INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABVIT415 standard; cDNA; 353 BP.
Human prostate expression marker cDNA 17406.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human prostate expression marker cDNA 47200
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         11.3%; Score 30.8; I
48.3%; Pred. No. 25;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              11.3%; Score 30.8; 50.0%; Pred. No. 29;
                                                                          ABX74144 standard; DNA; 32248 BP.
Human novel polynucleotide #972.
US2002132753-A1.
19-SEP-2002.
(ROSE) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABT09214 standard; DNA; 705 BP.
Phase-1 Rat CT gene SEQ ID No 302.
WO200266682-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADB12769 standard; DNA; 139308 BP.
Human PRKR DNA.
DE10128838-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABV47209 standard; cDNA; 450 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ACA22540 standard; DNA; 1236 BP
          (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                           11.4%;
                                                                                                                                                                                                    11.4%;
50.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        02-JAN-2003.
(GENP-) GENPROFILE AG.
                                                                                                                                                                                                                                                                                                       10-OCT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 414
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
                           Query Match
Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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Human musculoskeletal system related polynucleotide SEQ ID NO 3487.
WQ200155367-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Sed
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Human nervous system related polynucleotide SEQ ID NO 12743.
WO200159063-A2.
                                                                                                                                                                                                                                      ADE84149 standard; DNA; 4233 BP.
Human lymphoid cell proliferative disorder gene derived DNA
WO2003044226-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 32248
                                                                                                ADE84073 standard; DNA, 4233 BP. Human lymphoid cell proliferative disorder gene derived DNA WO2003044226-A2.
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                                                                                                                                                                                       Length 4233
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Human genomic DNA encoding partial novel secreted protein,
WO200155322-A2.
                                           Length 4233;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 5433;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABZ67645 standard; DNA; 8960 BP.
Human secreted protein encoding genomic DNA SEQ ID NO 1168
WO200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                              AAA99265 standard; DNA; 5433 BP.
Plasmodium yoelii YM MAEBL genomic DNA sequence SEQ ID
US6120770-A.
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N
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Human secreted protein encoding genomic DNA SEQ ID
WO200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 10;
63;
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63;
                                                                                                                                                                                                                                                                                   30-MAX-2003.
30-MAX-2003.
(EPIG-) EPIGENOMICS AG.
(EFY Match 11.4%; Score 31; DB 10;
(EFY Match 11.4%; Pred. No. 48;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN 11.4%; Score 31; DB 4;
ery Match 11.4%; Score 31; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                        USD. 2.2.000.
19-52P-2000.
(UNOT ) UNIV NOTRE DAME DU LAC.
(UNOT ) UNIV NOTRE DAME DU LAC.
(EXY MALCh 11.4%; Score 31; DB 3;
(EXY MALCh 53; Pred. No. 53;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ELY MATCh 11.4%; Score 31; DB 4;
ELY MATCh 50.3%; Pred. No. 1e+02;
                                                                                                                                                                                     Score 31; DB 10;
Pred. No. 48;
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8
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95;
                                               Score 31; DB
Pred. No. 48;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                03-0CT-2002.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) AUTCh
.ery Match
11.4%; Score 31;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      03-02-2002.
03-02T-2002.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 11.4%; Score 31;
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51.0%; Pred. No.
03-OCT-2002.
(EPIG-) EPIGENOMICS AG. 11.4%;
                                                                                                                                                                     AG.
11.4%;
49.1%;
                                                                                                                                                      30-MAY-2003.
(EPIG-) EPIGENOMICS
                                           Query Match
Best Local Similarity
RESULT 402
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 406
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Best Local Similarity
RESULT 408
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Best Local Similarity
RESULT 409
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RESULT 403
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Best Local Similarity
RESULT 404
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Best Local Similarity
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Query Match

RESULT

RESULT 405

PPRE

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(GENE-) GENE LOGIC INC.
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Best Local Similarity
REGULT 434
ID ADQ25068 standard; DN
DE Human soft tissue san
PN W02004048938-A2.
PD 10-UN-2004.
PA (PROT-) PROTEIN DESIG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 435
                                                                                                                                                                                                                                                                                                                           Best Local Similarity
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                                                                                                                                                                                                                               RESULT 429
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human immune/haematopoietic antigen genomic sequence SEQ ID NO:19535 WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAK75611 standard; DNA; 5314 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:30423
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAK91497 standard; DNA; 5314 BP.
Human digestive system antigen genomic sequence SEQ ID NO: 5073
WO200155314-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 30.8; DB 12; Length 76272; Pred. No. 1.6e+02;
                                                       11.3%; Score 30.8; DB 8; Length 1236; 54.4%; Pred. No. 36;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 2469
                                                                                                                                                                    11.3%; Score 30.8; DB 8; Length 1794; 51.4%; Pred. No. 41;
                                                                                                                                                                                                                                                                                   Length 1873;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 5314;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 5314;
                                                                                                                                                                                                                       DNA encoding novel human diagnostic protein #17144 WO200175067-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 30.8; DB 12;
Pred. No. 46;
                                                                                                                                                                                                                                                                                11.3%; Score 30.8; DB 5; 48.3%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Worderson.
09-AUG-2001.
(HTMA-) HUMAN GENOME SCI INC.
(HTMA-) HUMAN GENOME SCI INC.
11.3%; Score 30.8; DB 4;
ery Match
' nimilarity 51.4%; Pred. No. 60;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11.3%; Score 30.8; DB 4; 51.4%; Pred. No. 60;
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Oryza minuta Pi9 locus genomic DNA sequence.
US2004006788-A1.
U8-JAN-2004.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

(ery Match 11.3%; Score 30.8; I
                                                                                                             Prokaryotic essential gene #34310
W020027183-A2.
03-007-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                      ADJ40188 standard; cDNA; 2469 BP.
Plant cDNA #1188.
US2004016025-A1.
Prokaryotic essential gene #4197
W020027183-A2.
03-0021-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                               CDNA; 1873 BP
                                                                                                ACA52653 standard; DNA; 1794 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAK64723 standard; DNA; 5314 BP
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(HUMA-) HUMAN GENOME SCI INC.
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57.1%;
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52.3%;
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L-2001.

L-2Y MATCh

Best Local Similarity
RESULT 424
ID AAK64723 standa-
DE Human immur

PD 09-
PA
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JMA-) HUMAN GENOM.

JY MATCH

JEST LOCAL Similarity by

RESULT 426

ID AD157166 standar

DE OYZA minute

PN US20040067

PD 08-UAP
                                                                                                                                                                                                                                                                                                                                                                                                        BRIGGS S P.
COOPER B.
GLAZEBROOK J.
GOFF S A.
KATAGIRI F.
                                                                                                                                                                                                                                                                                                                                                           (BUDW/) BUDWORTH P. (MOUG/) MOUGHAMER T.
                                                                 Best Local Similarity RESULT 420
                                                                                                                                                                                Best Local Similarity
RESULT 421
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 425
                                                                                                                                                                                                                                                                                          Best Local Similarity
RESULT 422
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Best Local Similarity
                                                                                                                                                                                                            AAS81340 standard;
                                                                                                                                                                                                                                                     11-OCT-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (KREP/) KREPS J.
(PROV/) PROVART N.
(RICK/) RICKE D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ZHUT// ZHU T.
                                                        Query Match
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                                                                                                                                                                                                                                                                                                                                                                                           (MOUG/)
(BRIG/)
(COOP/)
(GLAZ/)
                                                                                                                                                                                                             PD PE
 PN
PD
PA
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Primary rat hepatocyte toxicity modelling related gene SEQ ID NO:898.
WO2003065993-A2.
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Tumour suppressor gene derived chemically modified sequence #460.
WO200168912-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 2377;
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(EPIG-) BPIGENOMICS AG.
11.2%; Score 30.6; DB 4; Length 6292;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB 4; Length 1587;
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                                                                                                                                 DB 10; Length 475
                                                                                                                                                                                                                                                                                                     Length 475
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 1000;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAH34864 standard; cDNA; 1587 BP.
Human colon cancer antigen encoding cDNA SEQ ID NO:1946
WO200122920-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MO2004048938-A2.
                                                                                                                                                                                                                                                                                                     DB 10;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DB 6;
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Query Match
Query Match
11.2%; Score 30.6; DB 1
Best Local Similarity 50.3%; Pred. No. 52;
RESULT 432
DB Haman secretory polynucleotide SPTM SEQ ID NO
PN WO200283876-A2.
ID ADB55776 standard; DNA; 475 BP.
DE TOxicity-related gene, SEQ ID 802.
PN WO2003064624-A2.
PD 07-AUG-2003.
PA (GENE-) GENE LOGIC INC.
Query Match
11.2%; Score 30.6; I Best Local Similarity 55.0%; Pred. No. 29;
                                                                                                                                                                                                                                                                                                                                                                                                                  (HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

(ery Match 11.2%; Score 30.6; I
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(HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

11.2%; Score 30.6; I

cery Match 52.8%; Pred. No. 45;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         WOLULUL
24-OCT-2002.
(INCY-) INCYTE GENOMICS INC.
(ELY MATCh 11.2%; Score 30.6; I
ELY MATCh 11.2%; Pred. No. 55;
                                                                                                                                                                                                                                                                                                Query Match 11.2%; Score 30.6; I
Best Local Similarity 55.0%; Pred. No. 29;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        11.2%; Score 30.6; 50.3%; Pred. No. 56;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          11.2%; Score 30.6; 50.3%; Pred. No. 62;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADB62455 standard; cDNA; 2942 BP.
Human cDNA encoding clone FEBRA20038330.
EP1308459-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADB62875 standard; cDNA; 2377 BP.
Human cDNA encoding clone OCBBF20177910.
BP1308459-A2.
07-MAY-2003.
                                                                                                                                                                                                                                                                                                                                                     ABL90179 standard; cDNA; 1000 BP.
Human polynucleotide SEQ ID NO 741.
WO200190304-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    07-MAY-2003.
(HELI-) HELIX RES INST.
(REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                            ADB50356 standard; DNA; 475 BP.
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ABZ71606 standard; cDNA; 1031 BP.
Breast specific nucleic acid # SEQ ID 66.
07.0020288375.A2.
07.NOV-2002.
(DIAD-) DIADEXUS INC.
                                                                                                                                                                                                                                                                                                                                                                                                          Prokaryotic essential gene #29681.
WO200277183-A2.
                                                                                                                                                                                                                                                                                    DNA; 1721 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADK14348 standard; DNA; 771 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 WOZUCZ-...
03-007-2002.
(ELIT-) ELITRA PHARM INC.
11.1%;
                                                                                     11.1%;
                                                                                                                                                                                                                               11.1%;
57.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Ouery Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 451
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 452
                                                                                                                                                                                                                                          Best Local Similarity
RESULT 446
                                                                                                                                                                                                                                                                                    ADL62875 standard;
                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
                                                                                                         Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                EP1033405-A2.
06-SEP-2000.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           23-AUG-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
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                                                                                         Query Match
                                                                                                                                                                                                                                 Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                                            ADI35046 standard; DNA; 322101 BP.
Human prostaglandin E receptor subtype EP3 (PTGER3) genomic DNA sequence.
US2003224393-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            אונים Standard; cDNA; 375 BP.
Bovine EST associated with lactation/muscle/fat deposition #15014.
US2002137139-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Bovine EST associated with lactation/muscle/fat deposition #3647 US2002137160-A1.
                                                                                                                                                                                                                                              07-AUG-2003.
(DECO-) DECODE GENETICS EHF.
ery Match 11.2%; Score 30.6; DB 10; Length 322101;
                                                                                                                                                                                                                                                                                                                                                                                                                           Length 322101
                                                                                                                                             Length 68571;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      2532
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          O2-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
HEY Match Human 11.1%; Score 30.4; DB 4; Length 612;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
11.1%; Score 30.4; DB 5; Length 619;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        11.1%; Score 30.4; DB 8; Length 324; 50.7%; Pred. No. 29;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 8; Length 375;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAL02531 standard; cDNA; 612 BP.
Human reproductive system related antigen cDNA SEQ ID NO:
W0200155320-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (HUMA.) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 57.3%; Pred. No. 37;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABL97297 standard; cDNA; 612 BP.
Human testicular antigen encoding cDNA SEQ ID NO:
W0200155317-A2.
                                                                                                                                                                                                                                                                                                                                                                                    Od-DEC-2003.
(DECO-) DECODE GENETICS EHF.
(DECO-) DECODE 11.2%; Score 30.6; DB 12;
ery Match 11.2%; Pred. No. 3e+02;
                                                                                                                                             11.2%; Score 30.6; DB 12; 53.8%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human prostate expression marker cDNA 53429.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.1%; Score 30.4; 148.8%; Pred. No. 31;
         60.0%; Pred. No. 74;
                                      ADH56913 standard; DNA; 68571 BP.

Human CARD4 genomic DNA sequence SeqID 1.
07S203219810-A1.
27-NOV-2003.
(BARN/) BARNES G.
(BERT/) BERIIN J.
                                                                                                                                                                                                  DNA; 322101 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABX53718 standard; cDNA; 324 BP
                                                                                                                                                                                                                 Human PAOD1 genomic DNA.
WO2003064471-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  26-SEP-2002.
(BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Lery Match
Best Local Similarity
RESULT 440
ID BAX49849
                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 438
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 441
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 442
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 443
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
                                                                                                                                                            Best Local Similarity RESULT 437
                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
RESULT 439
     Best Local Similarity RESULT 436
                                                                                                                                                                                                  AAD58431 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         02-AUG-2001
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
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Candida maltosa CYP gene upstream regulator region DNA SEQ ID NO:77. 26-FEB-2004. (COGN-) COGNIS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 26241;
                                 ADF85848 standard, CDNA, 1031 BP.
Human breast specific nucleic acid (BSNA) cDNA, SEQ ID NO:66.
W02003099989-A2.
04-DEC-2003.
%TO PADA.) DIADEXUS INC.
                                                                                                                                                                Length 1031;
                                                                                                                                                                                                                                                                                                                             Length 1721;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 771;
Length 1031;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 2538;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 490;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 3; Length 492;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 622
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABA16222 standard; DNA; 26241 BP.
Human nervous system related polynucleotide SEQ ID NO 8553
WO200159063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABQT6298 standard; cDNA; 896 BP.
S. cerevisiae BAX-associated cDNA fragment SEQ ID 21.
WO200264766-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AACS4789 standard; DNA, 490 BP.
Arabidopsis thallana DNA fragment SEQ ID NO: 79098.
BP1033405-A2.
06-SEP-2000.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAC38648 standard, DNA; 492 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 21740
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 12;
                                                                                                                                                                  DB 12;
                                                                                                                                                                                                                                                                                  27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              16-AUG-2001.

(HUMA-) HUMAN GENOME SCI INC.

Hery Match 11.1%; Score 30.4; DB 5;

Hery Match 59.1%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 3;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              s,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 8;
    DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. 
11.1%; Score 30.2; DB 
st Local Similarity 60.2%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABV55148 standard; cDNA; 622 BP,
Human prostate expression marker cDNA 55139-
WO200160860-A2.
    Score 30.4; I
Pred. No. 45;
                                                                                                                                                                  Score 30.4; I
Pred. No. 45;
                                                                                                                                                                                                                                                                                                                               11.1%; Score 30.4; I 52.3%; Pred. No. 54;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     11.1%; Score 30.2; 52.8%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11.1%; Score 30.2; 52.8%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          11.1%; Score 30.2; 51.9%; Pred. No. 46;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 30.4;
Pred. No. 62;
                                                                                                                                                                                                                                            Human ovarian cancer DNA marker #21087.
WO200170979-A2.
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Length 96598;

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AAD09356 standard; DNA; 1134 BP.
Human beta-1,3-galactosyltransferase homologue, ZNSSP8 degenerate DNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAH25483 standard; DNA; 180 BP.
Nuclectide fragment of ribulose-1,5-bisphosphate carboxylase cDNA.
WO200141559-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human immune/haematopoietic antigen encoding cDNA SEQ ID NO:8373. WO200157182-A2.
                                                                                                                                                  vuery Match
11.1%; Score 30.2; DB 10; Length 96598;
BBst Local Similarity 47.2%; Pred. No. 2.7e+02;
RESULT 464
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAX61837 standard; DNA; 1212 BP.
B. burgdorferi antigenic protein coding sequence, f04A.nt BB011.
WO9859071-A1.
                                                                                                                                                                                                                                                                                                                Score 30.2; DB 10; Length 96598;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                        (GAGR-) SAGRES DISCOVERY.
Query Match
11.1%; Score 30.2; DB 10;
Best Local Similarity 47.2%; Pred. No. 2.7e+02;
                    Score 30.2; DB 9;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 30; DB 4;
Pred. No. 32;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human leukocyte derived cDNA, 306 BP.
WO200257414-A2.
SP-ULL-2002.
(BIOC-) BIOCARDIA INC.
3=rY Match
11.0%; Score 30; DB 6; red. Incal Similarity 55.9%; Pred. No. 38.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 11.0%; Score 30;
ery match 55.9%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.0%; Score 30; 52.4%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              11.0%; Score 30; 23.8%; Pred. No.
                                                                                                                                                                                                                    ADE82941 standard; DNA; 96598 BP.
Mouse Nfkb1 gene genomic DNA sequence.
WO2003080808-A2.
                                                                                                                                                                                                                                                                                                                                                                 ADE95857 standard; DNA; 96598 BP.
Mouse Nfkb1 gene genomic DNA sequence.
WO2003039484-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         12-AUG-2003.
(GENO-) GENOME THERAPEUTICS CORP.
                                                                            DNA; 96598 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAK63313 standard; cDNA; 381 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADF02426 standard; DNA; 624 BP.
Bacterial polynucleotide #2711.
US6605709-B1.
                    11.18;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           11.0%;
                                                                                                                                                                                                                                                                                                              Y Match
Local Similarity 47.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   30-DEC-1998.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                          02-OCT-2003.
(SAGR-) SAGRES DISCOVERY.
 (SAGR-) SAGRES DISCOVERY
                                                                                                                             30-JAN-2003.
(SAGR-) SAGRES DISCOVERY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          21-JUN-2001.
(ZYMO) ZYMOGENETICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 14-JUN-2001.
(UNIC-) UNICROP LTD.
                                                                      ADB72347 standard;
Mouse Nfkbl gene.
WO2003008583-A2.
                    Query Match
Best Local Similarity
RESULT 463
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 468
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 469
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 470
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            WO200144479-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                            15-MAY-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Ouery Match
Best Local Si
RESULT 471
                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
                                                                                                                                                                                                                                                                                                                                                RESULT 465
                                                                                                                                                                                                                      ADH23603 standard; DNA; 1199 BP.
Baker's yeast small non-coding amplicon between H2a/H2b SeqID
W02003097868-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 10; Length 1308;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     11.1%; Score 30.2; DB 5; Length 15031; 51.9%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL17834 standard; DNA; 23203 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 4975.
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAL04361 standard; DNA; 32192 BP.
Human reproductive system related antigen DNA SEQ ID NO: 7049.
WO200155320-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 33053;
                                                                                                                                                                                                                                                                                                                                 Length 1199
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          11.1%; Score 30.2; DB 6; Length 1217; 58.2%; Pred. No. 55;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11.1%; Score 30.2; DB 4; Length 23203; 51.1%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 32192,
                                      DB 6; Length 896;
                                                                                                                                                                                    Length 987;
                                                                                                                                                                                                                                                                                                                                                                              AAS62692 standard; cDNA; 1217 BP. cDNA sequence #479 encoding novel human secreted protein WC200177291-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABG67005 standard; DNA; 33053 BP.
Human angiogenesis associated polynucleotide SEQ ID NO
WC200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADA02609 standard; DNA; 96598 BP.
Mouse NEkbl carcinoma associated gene, SEQ ID NO:1127.
17-JUL-2003.
17-JUL-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAS66875 standard; cDNA; 15031 BP.
DNA encoding novel human diagnostic protein #5679.
WO200175067-A2.
                                                                                                                                                                                                                                                                                                        PTY LTD.
11.1%; Score 30.2; DB 12;
49.1%; Pred. No. 54;
                                                                                                                                            17-DEC-1998.

(HUMA-) HUMAN GENOME SCI INC.
Query Match
Best Local Similarity 49.0%; Pred. No. 51,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.1%; Score 30.2; DB 4; 53.9%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 30.2; DB 6;
Pred. No. 1.8e+02;
                                                                                 Human secreted protein gene 21 clone HWTAZ75 WO9856804-A1.
                        uuery Match 11.1%; Score 30.2; I
Best Local Similarity 49.1%; Pred. No. 49;
RESULT 454
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             24-UN-2003,
(GENO-) GENOME THERAPEUTICS CORP.
ery Match 11.1%; Score 30.2; I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADC93856 standard; DNA; 1308 BP.
E. faecium DNA sequence SEQ ID 3483.
US6583275-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             11.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                  18-OCT-2001.
(GEMY) GENETICS INST INC.
                 (JANC ) JANSSEN PHARM NV.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     13-JUN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 488975 standard; cl
DD DNA encoding novel hr
PN WO20175067-A2.
PD 11-0CT-2001.
PA (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                         27-NOV-2003.
(MICR-) MICROBIOGEN
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                            Best Local Similarity
RESULT 456
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 457
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 459
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 460
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11-OCT-2001.
(HYSE-) HYSEQ INC.
22-AUG-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              27-SEP-2001
                                                                                                                                                                                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT 461
                                                                                                                                                                                                                                       A D D E C
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Length 1134;

DB 5; 62;

Length 624

DB 10; 50;

Length 381;

4,

DB 4

Length 306

9

Length 96598;

Length 180,

Length 5641;

Length 7432;

Length 7809

Length 8911;

Length 8911;

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ABZ10124 standard; DNA; 7809 BP.
Haematopoietic cell proliferation disorder related DNA sequence #264.
WO200277272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAK84636 standard; DNA; 8911 BP.
AAK84636 mtmune/haematopoietic antigen genomic sequence SEQ ID NO:39448.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human immune/haematopoietic antigen genomic sequence SEQ ID NO:39447.
WO200157182-A2.
           ABL21332 standard; DNA; 5641 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 15469.
W020011042-A2.
Z7-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         11.0%; Score 30; DB 1; Length 11873; 57.4%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 30; DB 8; Length 14067; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 30; DB 1; Length 11866; Pred. No. 1.4e+02;
                                                                                                                                         AAX56763 standard, DNA; 7432 BP.
Mouse odorant binding protein MMOBP1A genomic DNA.
DB19756678-C1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           11.0%; Score 30; DB 4; 1
55.9%; Pred. No. 1.3e+02;
                                                                                                      Score 30; DB 4; 1 Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                       Score 30; DB 8;
Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 30; DB 2;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                     Score 30; DB 2;
Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAX99556 standard; DNA; 14066 BP.
Nucleic acid sequence from U. urealyticum
WO9939007-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAN40141 standard; DNA; 11866 BP.
Sequence of human factor IX genomic DNA.
NO8400560-A.
16-FEB-1984.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAN40176 standard; DNA; 11873 BP.
Sequence of human factor IX genomic DNA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Prokaryotic essential gene #34468
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DNA; 14067 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DNA; 8911 BP
                                                                                                                                                                                                                                                                                                                                                    WO20us...
03-0CT-2002.
(EPIG-) EPIGENOMICS AG. 11.0%;
... Match ''arity 49.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            11.0%;
55.9%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        / Match 11.0%;
Local Similarity 57.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            11.0%;
                                                                                                      11.0%;
                                                                                                                                                                                                                                                       11.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (NATR ) NAT RES DEV CORP. (BROW/) BROWNLEE G G.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    07-MAR-1984.
(NATR ) NAT RES DEV CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          05-AUG-1999.
(UABR-) UAB RES FOUND.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 487
                                                                                                                                                                                                                   20-MAY-1999.
(FORS/) FORSSMANN W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 488
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 482
                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACA52811 standard;
                                                                                                                           Local Similarity
                                                                                                                                         RESULT 481
ID AAX56763 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAK84635 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         GB2125409-A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
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                                                                                                          Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                              Best Loca
RESULT 483
RESULT 480
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAN91624 standard; DNA; 2160 BP.
5' flanking sequence of sigma-gamma-4 switch sequence for Cx gene
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         NO 15472
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.0%; Score 30; DB 12; Length 3741; 55.9%; Pred. No. 95;
                                                                                                                                                                                                                                                                                                                 Length 1590;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.0%; Score 30; DB 6; Length 1917; 54.5%; Pred. No. 74;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (37-JAN-2003.
(37-JAN-2003.
(37GN ) SYNGENTA PARTICIPATIONS AG.
ery Match ** Match ** 11.0%; Score 30; DB 8; Length 2000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 2160;
                 DB 2; Length 1212;
63;
                                                                                                                                                                DB 8; Length 1575;
69;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11.0%; Score 30; DB 3; Length 1671; 50.7%; Pred. No. 71;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 3144
                                                                                                                                                                                                                                                                                                                                                                      AAC76388 standard; cDNA; 1671 BP.
Human ORFX ORF1943 polynucleotide sequence SEQ ID NO:3885
WO200058473-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADQ22350 standard; DNA; 3741 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 5170.
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABK75003 standard; DNA; 1917 BP.
Bacillus licheniformis genomic sequence tag (GST) #2294.
W0200229113-A2.
11-ARP-2002.
(NOVO ) NOVOZYMES BIOTECH INC.
(NOVO ) NOVOZYMES AS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL21333 standard; DNA; 3144 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID.
WO200171042-A2.
                                                                                                                                                                                                                                                                       12-003.

12-006-2003.

(GENO-) GENOME THERAPEUTICS CORP.

6IV MAtch 11.0%; Score 30; DB 10;

ery Match 59.3%; Pred. No. 70;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 4;
89;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 1;
78;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.0%; Score 30; 53.4%; Pred. No.
             Query Match 11.0%; Score 30;
Best Local Similarity 45.7%; Pred. No.
RESULT 472
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11.0%; Score 30; 52.4%; Pred. No.
                                                                                                                                                                  Score 30;
Pred. No.
                                                                                       Prokaryotic essential gene #26261.
W0200277183-A2.
03-OCT-2002.
(ELITRA PHARM INC.
sry Match
EL Local Similarity 59.3%; Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADA73195 standard; DNA; 2000 BP.
Rice gene, SEQ ID 6521.
WO2003000898-A1.
                                                                                                                                                                                                                        DNA; 1590 BP.
                                                                       ACA44604 standard; DNA; 1575 BP
                                                                                                                                                                                                                                        Bacterial polynucleotide #2561.
US6605709-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 20-APR-1989.
(MITK) MITSUI TOATSU CHEM INC.
(MITH) MISTUI PHARM INC.
(EDUC-) EDUCATION FOUND FUJITA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (FUJI-) FUJITA GAKYEN GH. (FUJI-) FUJITA GAKYEN GH.
(MEDI-) MEDIMMUNE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                              05-OCT-2000.
(CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Ouery Match
Best Local Similarity
RESULT 478
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 476
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 479
                                                                                                                                                                                  Best Local Similarity RESULT 473
                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 474
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 475
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
                                                                                                                                                                                                                          ADF02276 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                upstream.
WO8903425-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
                                                                                                                                                                      Query Match
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RESULT 477

ID DE PN PD

Length 14066;

Length 61457;

Length 67191;

```
11.0%; Score 30; DB 11; Length 67191; 61.3%; Pred. No. 2.7e+02;
                                                                                                                                                               Score 30; DB 9; Length 67191;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Human soft tissue sarcoma-upregulated DNA - SEQ ID 3426
WO2004048938-A2.
                                                                    ADA02903 standard; DNA; 67191 BP.
Mouse Mcm3ap carcinoma associated gene, SEQ ID NO:1421.
17-JUL-2003:
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ACF68988 standard; DNA; 681 BP.
Photorhabdus luminescens nucleotide sequence #7455.
WO200294867-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADM74498 standard; DNA; 67191 BP.
Wurine carcinoma associated (CA) nucleic acid #85.
WS2004072154-A1.
15-APR-2004.
                                                                                                                                                                                                                                                                                                          Score 30; DB 10;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 30; DB 12;
Pred. No. 2.7e+02;
                      Score 30; DB 10;
Pred. No. 2.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 30; DB 2;
Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
ery Match
ft Local Similarity 46.6%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.9%; Score 29.8; 1 52.9%; Pred. No. 53;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.9%; Score 29.8; 163.0%; Pred. No. 57;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ACA39303 standard; DNA; 614 BP.
Prokaryotic essential gene #20960.
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADQ20606 standard; DNA; 195917 BP
                                                                                                                                                                                                                 ADB72641 standard; DNA; 67191 BP
                                                                                                                                                                                                                                                                                                                                                             DNA; 67191 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACH22153 standard; cDNA, 498 BP.
Human adult ovary cDNA #533.
US2003073623-Al.
                                                                                                                                                               11.0%;
61.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.0%;
57.4%;
                    11.0%;
                                                                                                                                                                                                                                                                                                         11.0%;
61.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                11.0%;
61.3%;
                                                                                                                                                                                                                                                                                                                                                           ADL95495 standard; DNA; 67
Murine MCM3AP genomic DNA.
US2003165878-A1.
   (SAGR-) SAGRES DISCOVERY
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                        Query Match 11.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PA (LABA/) LABAT I.

PA (STAC/) STACHE-CRAIN B.

PA (JOKE/) DICKSON M.C.

QUETY MATCH
10.9
                                                                                                                                                                                                                                                                                                                                                                                                             04-SEP-2003.
(MORR/) MORRIS D W.
(ENGE/) ENGELHARD E K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (MORR/) MORRIS D W. (ENGE/) ENGELHARD E K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    17-APR-2003.
(DRMA/) DRMANAC R T.
                    Query Match
Best Local Similarity
RESULT 498
                                                                                                                                                                       Best Local Similarity
RESULT 499
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
RESULT 502
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 503
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
RESULT 505
                                                                                                                                                                                                                                                                                                                         Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Local Similarity
                                                                                                                                                                                                                                      Mouse Mcm3ap gene. WO2003008583-A2.
                                                                                                                                                                                                                                                                       30-JAN-2003
                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                          AAK89418 standard; DNA; 28588 BP.
Human digestive system antigen genomic sequence SEQ ID NO: 2994
WO200155314-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 38059;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 28588;
                                                                                                                            Length 18359;
                                                                                                                                                                                                                         (HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

(Lery Match 11.0%; Score 30; DB 4; Length 28588; Lery Match 11.0%; Fred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            11.0%; Score 30; DB 4; Length 38059; 57.4%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                               Length 28588;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           11.0%; Score 30; DB 6; Length 28588; 61.5%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABN90278 standard; DNA; 28588 BP.
Human liver antigen HLDRI94 genomic sequence, SEQ ID NO:399.
US2002042096-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 38059;
                AAX20255 standard; DNA; 18359 BP.
Borrelia burgdorferi polynucleotide sequence #8.
WO9858943-A1.
                                                                                                                   Query Match
Best Local Similarity 45.7%; Pred. No. 1.7e+02;
RESULT 490
DE AAK89418 standard; DNA; 28588 BP.
DB Human digestive system antigen genomic sequence PD 0.2-40(2-2011.
PD 0.2-40(2-2011.
PA (HUMA-) HUMAN GENOME SCI INC.
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24-APR-2003.
(HUMA.) HUMAN GENOME SCI INC.
(HIMA.) HUMAN GENOME SCI INC.
11.0%; Score 30; DB 11;
ery Match
11.0%; Score 30; DB 11;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            TI-ARE-SCOURT (GENE LOGIC INC. 11.0%; Score 30; DB 6; 7 Y Match 11.0%; Pred. No. 2.2e+02; Frocal Similarity 57.4%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   11.0%; Score 30; DB 10; 57.4%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADD71098 standard; DNA; 38059 BP.
Human coagulation factor IX gene SEQ ID NO:102.
WO2003061564-A2.
                                                                                                                                                                                                                                                                                                                                                                                                           11.0%; Score 30; DB 5; 61.5%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human liver-related genomic DNA - SEQ ID 399 US2003077602-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAF54018 standard, DNA, 38059 BP.
Human factor IX (hFIX) gene, SEQ ID NO:4.
14-DEC-2000.
(UNMI ) UNIV MICHIGAN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABN95627 standard; DNA; 38059 BP.
Gene #2125 used to diagnose liver cancer.
WO200229103-A2.
                                                                                                                                                                                                                                                                                                                    AAS31923 standard, DNA; 28588 BP.
Human liver associated genomic DNA #97.
WO200155355-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADJ15191 standard; DNA; 28588 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADC85383 standard; DNA; 61457 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Mouse Mcm3ap genomic sequence. WO2003045230-A2. 05-JUN-2003.
                                                                                                                                                                                                                                                                                                                                                                        02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                               30-DEC-1998.
(HUMA-) HUMAN GENOME SCI INC.
(MEDI-) MEDIMMUNE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (GENE-) GENE LOGIC INC. (LGBI-) LG BIOMEDICAL INST.
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J-A2.

J-2000.

Lery Match

Best Local Similarity 5
RESULT 495
ID ABN95627 standa-
DE Gene #2125
PD WQ200227
PD 11-**
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               11-APR-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                               Best Local Similarity
RESULT 491
                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
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RESULT 496
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Best Local Similarity
RESULT 493
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RESULT 489
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Length 67191;

Length 110000;

Length 195917;

DB 9; Length 498;

Length 614;

8;

DB

Length 6303;

Length 6352;

Length 6352;

Query Match

Query Match

Length 6352;

Length 7589

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Human musculoskeletal system related polynucleotide SEQ ID NO 3446.
WO200155367-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human musculoskeletal system-associated genomic DNA - SEQ ID 3446 US2004009488-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABX60069 standard; cDNA; 13605 BP. cDNA encoding novel human musculoskeletal system antigen #2413. US2002147140-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                  AAS61235 standard; DNA; 6352 BP.
Human gene regulation-associated gene oligonucleotide
WO200177375-A2.
                                                                                                                                                                                                                                                  ABL70563 standard; DNA; 6352 BP.
Chemically treated cell signalling DNA sequence#227
WO200202807-A2.
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15-JAN-2004.

15-JAN-2004.

HUMA-) HUMAN GENOME SCI INC.

(ery Match 10.9%; Score 29.8; DB 12;

(ery Match 10.9%; Score 29.8; DB 12;

(ery Match 10.9%; Score 29.8; DB 12;
                                                                                               ABK31340 standard; DNA; 6352 BP.
Signal transduction associated gene modified DNA
WO200200926-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           WOACCEDON.
02-AUG-2001.
(HTMA-) HUMAN GENOME SCI INC.
(HTMA-) HUMAN GENOME SCI INC.
10.9%; Score 29.8; DB 4;
ery Match
10.9%; Pred. No. 1.7e+02;
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(EPIG-) BPIGENOMICS AG.
(ery Match 10.9%; Score 29.8; DB 6;
ery Match 1.4e+02;
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Pred. No. 1.9e+02;
                                     10.9%; Score 29.8; DB 4; 51.9%; Pred. No. 1.3e+02;
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Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 29.8; DB 6;
Pred. No. 1.3e+02;
                                                                                                                                                                                                Score 29.8; DB 6;
Pred. No. 1.3e+02;
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Pred. No. 1.3e+02;
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Human immune system associated gene SEQ ID NO: W020020928-A2.
03-JAN-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DNA transcription associated genomic DNA #132.
WO200192565-A2.
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Human chemically treated genomic DNA #24
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DNA; 13605 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  BP.
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18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
10.9%;
... Match ... natch ... 88.4%;
09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                           WOZUGECCO
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.9%;
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51.1%;
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                                                                                                                                                                                                                                                                                                                   PD 10-JAN-2002.

PA (EPIG-) EPIGENOMICS AG.

Query Match 10.9%;

Best Local Similarity 58.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (EPIG-) EPIGENOMICS AG.
ry Match 10.9%;
t Local Similarity 51.9%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10-OCT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                 Best_Local_Similarity
RESULT 516
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
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Best Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABK28389 standard;
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Human immune/haematopoietic antigen genomic sequence SEQ ID NO:20945.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human immune/haematopoietic antigen genomic sequence SEQ ID NO:20946.
WO200157182-A2.
09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL26452 standard; DNA; 5646 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 30829
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                Length 1098;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 2817;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 6295;
                                 10.9%; Score 29.8; DB 10; Length 681; 49.7%; Pred. No. 59;
                                                                                                                                                                                                     Length 690;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 5; Length 2817;
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                                                                                                                                                                                                                                                                                   #24
                                                                                                                                                                                                                                                                                                                                                                                                      AAS//233 standard; cDNA; 2817 BP.
DNA encoding novel human diagnostic protein #13037.
W2200175067-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       cDNA; 5962 BP.
human diagnostic protein #14938
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  cDNA; 2817 BP.
human diagnostic protein #14433
                                                                                                                                                                                                                                                         ADK16096 standard; DNA; 1098 BP.
Nanoarchaeum equitans cancer-associated (CA) gene WO2003093434-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.9%; Score 29.8; DB 10; 49.1%; Pred. No. 1.1e+02;
                                                                                                                                                                                                     DB 12;
                                                                                                                                                                                                                                                                                                                                                                DB 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.9%; Score 29.8; DB 5; 60.5%; Pred. No. 1.3e+02;
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  PA (INSP ) INST PASTEUR.

PA (CMRS) CARS CENT NAT RECH SCI.

Query Match

Best Local Similarity 49.7%; Pred. No. 59;

RESULT 507

RESULT 507

DE DNA encoding a M. catarrhalis protein #1512.

PD (JAN-2004.

PA (GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.9%; Score 29.8; DB 60.5%; Pred. No. 99;
                                                                                                                                                                                                   10.9%; Score 29.8; 1
63.0%; Pred. No. 59;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.9%; Score 29.8; 1 60.5%; Pred. No. 99;
                                                                                                                                                                                                                                                                                                                                                              10.9%; Score 29.8; 52.9%; Pred. No. 70;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human cDNA encoding clone BRSSN20001970 EP1308459-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADB62147 standard; cDNA; 3685 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (HELL-) HELIX RES INST.
(REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAK66134 standard; DNA; 6295 BP.
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RESULT 5.1

ID ADBG5147 standard; cI
DE Human cDNA encoding c
PN EP1308459-A2.
PD 07-MXY-2003.
PA (RELL-) RES ASSOC BIC
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Best Local Similarity
RESULT 513
ID AAS/9134 standard; cl
DB DNA encoding novel hr
PN WO200175067-A2.
                                                                                                                                                                                                                                                                                                                                            (DIVE-) DIVERSA CORP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAS78629 standard; c
DNA encoding novel h
WOZ00175067-A2.
11-OCT-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
RESULT 515
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Best Local Similarity
RESULT 512
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RESULT 508
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RESULT 509
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Best Local Similarity
RESULT 510
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(PEKE ) PE CORP NY
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(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11-OCT-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          WO200157182-A2.
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Length 13605;

Match

Length 13605;

Length 13605;

Length 16373;

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17-APR-2003
                                                                                                 Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.9%; Score 29.8; DB 10; Length 110000; 49.7%; Pred. No. 3.7e+02;
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Best Local Similarity 51.1%; Pred. No. 4.18+02;
BESULT 531

ID ABZ72040 standard; DNA; 207433 BP.

BE Gene 216 H194BAC1098L22 nucleotide sequence SEQ ID NO 5.

PD 25-CCT-2001.

PA (GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                                                                                                                                                                                                                                                             10.9%; Score 29.8; DB 12; Length 76341; 60.5%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                RESULT 530

ID ABK83574 standard; cDNA; 147419 BP.

WO200228999-A2.

PD 11-APR-2002.

PD 11-APR-2002.

PAR-2002.

PAR-2002.

PAR-2002.

PAR-2002.

PAR-2002.

PAR-2002.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.9%; Score 29.8; DB 8; Length 207433; 55.2%; Pred. No. 4.6e+02;
                                                                                                                                                                                                              Length 62909;
                                                           Length 16373;
                                                                                                                                                                                                                                                                                   Prostate and breast cancer associated human gene SRD5A2 w02040208346-A2.
08-APR-2004 (AMSH ) AMERSHAM BIOSCIENCES SV CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match 10.9%; Score 29.8; DB 12;
Best Local Similarity 52.9%; Pred. No. 3.7e+02;
                                                                                                                                                    PD 28-DEC-2000.

PA (INCY-) INCYTE GENOMICS INC.

Query Match

Best Local Similarity 63.0%; Pred. No. 3e+02;

RESULT 526
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Oct. Co. Co. Co. California. (REC) UNIV CALIFORNIA. (REC) UNIV CALIFORNIA. 10.9%; Score 29.8; DB 8; ery Match 10.9%; Pred. No. 3.7e+02;
                                                       10.9%; Score 29.8; DB 6; 51.9%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADJ36614 standard; DNA; 207433 BP.
Bacterial artificial chromosome RPCI-11
US2004002470-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                  ABX16390 standard; DNA; 659158 BP. Mouse high growth region. US2002155564-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADK16049 standard; DNA; 490885 BP. Nanoarcheeum equitans genome. WOZO03093434-A2. 13-NOV-2003. (DIVE-) DIVERSA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AEX74891 standard; DNA; 207433 BP. BAC1098122 DNA sequence. W0200283077-A2. 24-OCT-2002. (SCHE ) SCHERING CORP. (GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           25-OCT-2001.
(GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                 DNA; 62909 BP
                                                                                                                                                                                                                                                                 ADM97422 standard; DNA; 76341 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (KEIT/) KEITH T.
(LITT/) LITTLE R D.
(VEER/) VAN EERDEWEGH P.
                 10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       11-APR-2002.
(GENE-) GENE LOGIC INC.
                                                                                                         AAF28545 standard; DN
Genomic fragment #32.
WO200078968-A2.
                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 527
                                                     Query Match
Best Local Similarity
RESULT 525
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 529
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 528
WO200202809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         RESULT 533
ID ADJ366
DE Bacter
PN US2004
PA (KEIT/
PA (LITT/
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Score 29.8; DB 12; Length 207433; Pred. No. 4.6e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 12; Length 207433;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 3; Length 1193;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 6; Length 352;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 476;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 6; Length 511;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human ovarian antigen HODHE36 CDNA, SEQ ID NO:1497.

NO200200677-Al.

O3-JAN-2002.

A (HUMA-) HUMAN GENOME SCI INC.

Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Photorhabdus luminescens nucleotide sequence #33
WO200294867-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 29.8; DB 12
Pred. No. 4.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WALCOURT OF THE STATE OF STATE O
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.8%; Score 29.6; I
51.5%; Pred. No. 54;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.8%; Score 29.6; 50.0%; Pred. No. 60;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.8%; Score 29.6; 148.7%; Pred. No. 62;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AACS9912 standard; cDNA; 1193 BP.
Human secreted protein cDNA sequence #6.
WO200055198-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADL63683 standard; DNA; 1330 BP.
Human ovarian cancer DNA marker #21895.
WO200170979-A2.
27-SEP-2001.
                                                                                                                                                                                                                                                                                 RESULT 534
ID ADL81193 standard; DNA; 207433 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ACF65380 standard; DNA; 210710 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ACH41552 standard; cDNA; 476 BP.
Human foetal brain cDNA #2919.
US2003073623-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity 50.0%; Pre
RESULT 538
ID ABQ55617 standard; cDNA; 511 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             28-NOV-2002.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                         10.9%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.9%;
55.2%;
(DUPU/) DUPUIS J.
(DMAS/) DEE MASTRO R G.
(SINO/) SIMON J.
(ALLE)/ ALLEN K.
(PAND/) PANDIT S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (KEIT) KEITH T.

LITT/) LITTLE R D.

(EERD/) ERRDEWEGH P V.

(DUDV/) DUPULS J.

(DWAS/) DEL MASTRO R G.

(SIMO/) SIMON J.

(ALLE/) ALLEN K.

(PAND/) PANDIT S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (DRWA/) DRWANAC R T.
(LABA/) LABAT I.
(STAC/) STACHE-CRAIN B.
(DICK/) DICKSON M C.
(JONE/) JONES L W.
                                                                                                                                                                                                                                                                                                                                                              BAC1098L22 DNA sequence
US2004023215-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    21-FEB-2002,
(CHIR ) CHIRON CORP.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 535
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RESULT 539
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
                                                                                                                                                                                                                                         Local Similarity
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ABL28044 standard; DNA; 14771 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 35605.
WO200171042-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 14771;
                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 6074;
   Length 3865;
                                                                                                                                                                                                                                                                                                                Length 5982;
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                                                                                                                                                  Length 4333;
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WO2002103041-A2.
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                                                                             NO:1
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Human immune system associated gene SEQ ID NO: 1653.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human angiogenesis associated polynucleotide SEQ WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AG.
10.8%; Score 29.6; DB 6;
56.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 29.6; DB 8;
Pred. No. 1.5e+02;
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Pred. No. 1.5e+02;
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Pred. No. 2.1e+02;
                                                                                                           27-MAR-2003.
(USSH ) US DEPT HEALTH & HUMAN SERVICES.
ery Match
10.8%; Score 29.6; DB 8;
ery Match 57.6%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 29.6; DB 8;
Pred. No. 1.5e+02;
     10.8%; Score 29.6; DB 2; 54.6%; Pred. No. 1.3e+02;
                                                                         Human transductin-1 (TDC1) encoding cDNA SEQ ID WO2003025140-A2.
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Human metastasis associated gene SEQ ID NO: 133
                                                                                                                                                                                                                                                                                                              10.8%; Score 29.6; DB 6; 53.4%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 29.6; DB 6;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 sample #22
                                                                                                                                                                                                                                                                                                                                                                                  Human DNA for staging of Astrocytomas #38
WO200202808-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADA20378 standard; DNA; 6074 BP.
Prostate tumour related genomic DNA sa W02002103042-A2.
27-DEC-2002.
(EPDG-) EPIGENOMICS AG.
20ery Match
10.8%; Score 29.
Sext Local Similarity 54.6%; Pred. No.
                                                                                                                                                                                          Listeria innocua DNA; 5982 BP.
WO20022891-A2,
                                                            CDNA; 4333 BP.
                                                                                                                                                                                                                                                                                                                                                                     ABK33990 standard; DNA; 6074 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             BP
                                                                                                                                                                                                                                                                            (INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI
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13-JUN-2002
(EPIG-) EPIGENOMICS AG.
10.8%;
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(EPIG-) EPIGENOMICS AG.
10.8%;
10.70.21 Similarity 54.6%;
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56.0%;
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27-DEC-2002
(EPIG-) EPIGENOMICS AG.
10.8%;
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(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 557
Query Match
Best Local Similarity
RESULT 550
                                                                                                                                                                                                                                                                                                                                 Local Similarity
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Best Local Similarity
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(PEKE ) PE CORP NY.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADA84185 standard;
                                                            ACC69613 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 WO200177376-A2.
18-OCT-2001.
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RESULT 556
                                                                                                                                                                                                           PA PRE PA
                                                                                                                                                                                                         AAH52576 standard; DNA; 1599 BP.
S. epidermidis open reading frame nucleotide sequence SEQ ID NO:545.
W0200134809-A2.
17-MAY-2001.
(CLAX.) GLAXO GROUP LTD.
                                                                                                                                                                                                                                                                                                                                                                       ABN90872 standard; DNA; 1692 BP. Staphylococcus epidermidis ORF nucleic acid sequence SEQ ID NO:335. USG380370-B1. 30-APR-2002.
                                                                   ABN91152 standard; DNA; 1404 BP.
Staphylococcus epidermidis ORF nucleic acid sequence SEQ ID NO:615.
US6380370-B1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAH55029 standard, DNA, 3398 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:4393
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   S. epidermidis genomic polynucleotide sequence SEQ ID NO:3936 WO200134809-A2.
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(GLAX ) GLAXO GROUP LTD.
ery Match 10.8%; Score 29.6; DB 4; Length 3609;
ery Match 52.4%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            29.6; DB 8; Length 1701;
No. 95;
                                                                                                                                                                                                                                                                                                                                                                                                                                                               DB 6; Length 1692;
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                            Length 1330;
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                                                                                                                                                                         Length 1404
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Human angiogenesis associated polynucleotide SEQ ID NO
WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        64
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Intron 3 of human pRb2/pl30 tumour suppressor gene
WO9738125-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Π
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (FIVE-) FIVE PRIME THERAPEUTICS INC.

10.8%; Score 29.6; DB 12;

t Local Similarity 53.4%; Pred. No. 1.1e+02;
                                                                                                                             30-ASR-2002.
(GENO-) GENOME THERAPEUTICS CORP.
(ery Match 10.8%; Score 29.6; DB 6;
         (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. ry Match 10.8%; Score 29.6; DB 5; t Local Similarity 54.6%; Pred. No. 87;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 29.6; DB 6;
Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 29.6; DB 4;
Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                    DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ACC62228 standard; cDNA; 1701 BP.
Human secreted protein #54 coding sequence SEQ
WO200299066-A2.
                                                                                                                                                                                                                                                                                                                    10.8%; Score 29.6; 1 52.4%; Pred. No. 93;
                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.8%; Score 29.6; 52.4%; Pred. No. 95;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.8%; Score 54.6%; Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                              (GENO-) GENOME THERAPEUTICS CORP. ry Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AD035510 standard; DNA; 2712 BP.
Novel mouse gene sequence #183.
W02004046310-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAH54572 standard; DNA; 3609 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        16-OCT-1997.
(UYJE-) UNIV JEFFERSON THOMAS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.8%;
54.6%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
RESULT 548
ID AAH54572 standard; D
B S. epidermidis genom
PW W0200134809-A2.
PD 17-MAY-2001.
PAAY | GLAX |
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 549
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 544
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Best Local Similarity
RESULT 546
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Best Local Similarity
RESULT 547
                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 542
                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 543
                              Query Match
Best Local Similarity
RESULT 541
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Local Similarity
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                                                                                                                                                                                                                                                                                                                    Query Match
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RESULT 545
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ID DE PN PD

PNED

P P P P P

DB 10; Length 1191;

Length 1029;

DB 8;

Length 1342;

ID DE PN PD

Length 2555;

Length 3095;

Length 6167;

615.

Length 6167;

Length 6167;

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AAS45136 standard; DNA; 6167 BP. Chemically pretreated genomic DNA associated with cell cycle #21. WO200168911-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Drosophila melanogaster genomic polynucleotide SEQ ID NO 29113
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
ery Match 10.8%; Score 29.4; DB 4; Length 6167;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human gene regulation-associated gene oligonucleotide #106.
                                                                                                                                        ABK17910 standard; CDNA; 1191 BP.
CDNA encoding human G-protein coupled receptor GCREC-64
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human cDNA differentially expressed in lung cancer #510 US2003065157-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABL70193 standard; DNA; 6167 BP. Chemically treated cell signalling DNA sequence#42.WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                            (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
(ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
ery Match
10.8%; Score 29.4; DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 29.4; DB 4;
Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 29.4; DB 6;
Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            29.4; DB 9;
No. 1.4e+02;
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Pred. No. 1.8e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABL32642 standard; DNA; 6167 BP.
Human immune system associated gene SEQ ID NO:
WO20020928-A2.
03-JAN-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABK28171 standard; DNA; 6167 BP.
DNA transcription associated genomic DNA #23.
W0200192565-A2.
(EPIG-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                             PA (INCY-) INCYTE GENOMICS INC.
Query Match
10.8%; Score 29.4; I
Best Local Similarity 60.8%; Pred. No. 97;
RESULT 571
                                                                                       Score 29.4; | Pred. No. 92;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score
Pred.
                                                                                                                                                                                                                                                                                                                                                                        ADC86322 standard; DNA; 1342 BP.
Human GPCR gene SEQ ID NO:775.
EP1270724-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACH04305 standard; cDNA; 3095 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABL25880 standard; DNA; 2555 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAS61151 standard; DNA; 6167 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.8%;
51.1%;
                                                                                       10.8%;
48.0%;
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4 (EPIG-) BPIGENOMICS AG.

10-EY MATCH

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(BPIG-) EPIGENOMICS AG. 10.8%;
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                                   03-OCT-2002.
(ELIT-) ELITRA PHARM INC
                                                                                                    Best Local Similarity RESULT 570
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 572
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WO200171042-A2.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 575
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
RESULT 573
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 574
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(LASE/) LASEK A W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Local Similarity
             WO200277183-A2.
                                                                                                                                                                                                                                          10-OCT-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                       Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AASO9301 standard; DNA; 319608 BP.
Human schizophrenia associated gene g35030 and biallelic markers Al-A71.
ery Match
st Local Similarity 54.6%; Pred. No. 6.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Haemophilus influenzae DNA for cellular proliferation protein #6 WO200170955-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL28042 standard; DNA; 19674 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 35599.
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 110000;
     Score 29.6; DB 6; Length 15951; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 110000,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 134499;
                                                                                                                                                                                                               Length 15951;
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                                                                                                                                                                                                                                                                                                                                                                                                                              Length 19634;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 19674;
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                                                                                                                                                                                                                                                                                   ABZ09891 standard; DNA; 19634 BP.
Human 5' and/or regulatory region of GPR37 DNA SEQ ID NO:31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAH51601 standard; DNA; 319608 BP.
Human chromosome 13q31-q33 genomic nucleotide sequence
WO200058510-A2.
                                                                         ABL70373 standard, DNA; 15951 BP.
Chemically treated cell signalling DNA sequence#132.
W02002807-A2.
10-JAN-2002.
(EPIG-) BPIGENOMICS AG.
ery Match
10.8%; Score 29.6; DB 6; Lerst Match
55.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                   Abc.

Human 5, anu,

WO200277272-A2.

03-OCT-2002.

(EPIG-) EPIGENOMICS AG.

Match 10.8%; Score 29.6; DB 8;

Match 7.**ity 54.6%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 29.6; DB 6;
Pred. No. 4.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    11-APP-2002.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
(CNRS ) CNRS CENT NAT 8CH SCI.
(CNRS ) CNRS CENT NAT 8CH SCI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 29.6; DB 4;
Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.8%; Score 29.6; DB 3; 51.5%; Pred. No. 4.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 29.6; DB 6;
Pred. No. 4.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 29.6; DB 3;
Pred. No. 6.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAF22286 standard; DNA; 134499 BP.
BAC containing repeats from centromeres 1-4 #9.
WO200055325-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 29.4; I
Pred. No. 92;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AEQ67197 standard; DNA; 1163020 BP.
Listeria innocua contig DNA sequence #10.
WO200228891-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ACA34007 standard; DNA; 1029 BP.
Prokaryotic essential gene #15664
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAS53224 standard; DNA; 1029 BP
     10.8%;
56.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.8%;
53.4%;
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54.6%;
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48.0%;
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53.4%;
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(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 INST PASTEL

S.S.) CNRS CENT NA.

Best Local Similarity 5:

RESULT 565

ID AAF22286 standa-

DE BAC contain

PN WO20005-

PA 21-7

PA
Query Match
Best Local Similarity
REGULT 559
ID ABL/0373 standard; Di
DE Chemically treated co
PN W0200202807-A2.
PD 10-DAN-2002.
PA (EPIG-) EPIGENOMICS;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 568
ID AAS5324 standard; DI
DE Haemophilus influenz;
PN WO20170955-A2.
PD 27-SEP-2001.
PA (ELIT-) ELITRA PHARM
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Best Local Similarity
RESULT 566
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Best Local Similarity
RESULT 569
                                                                                                                                                                                                         Query Match
Best Local Similarity
RESULT 560
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Best Local Similarity
RESULT 564
                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 561
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 562
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best_Local Similarity RESULT 567
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 563
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   05-OCT-2000.
(GEST ) GENSET.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
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PA

Length 42979

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AAD07868 standard; cDNA; 820 BP.
Human secreted protein-encoding gene 13 cDNA clone HIBEB47, SEQ ID NO:70.
W0200132675-A1.
                                                                                             Human corneal N-acetylglucosamine-6-sulfotransferase DNA SEQ ID NO:38.
US2002061562-Al.
(FUKU) FUKUDA M N.
(AKM) AKAMA T O.

YA MARCH
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADB50217 standard; DNA; 588 BP.
Primary rat hepatocyte toxicity modelling related gene SEQ ID NO:759.
WO2003065993-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABQ83210 standard; DNA; 397658 BP.
Human transporter protein genomic DNA SEQ ID NO:3.
ery Match
10.8%; Score 29.4; DB 8; Length 110000;
st Local Similarity 55.3%; Pred. No. 4.9e+02;
ABL20870 standard; DNA; 42979 BP. Drosophila melanogaster genomic polynucleotide SEQ ID NO 14083 WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABIT44365 standard; DNA; 113000 BP.
Partial genomic sequence of human oestrogen receptor beta DNA W02003050133-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Nuclectide sequence of a fragment of Salmon fish DNA. 19-202101090-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 29.4; DB 9;
Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                .,
                                                                                                                                                                                                                                                                                                                                                                                                                  10.8%; Score 29.4; DB 2; 48.0%; Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.8%; Score 29.4; DB 4; 49.7%; Pred. No. 7.2e+02;
                                                                                                                                                                                                                                                   Score 29.4; DB 6;
Pred. No. 3.7e+02;
                                                                                       Score 29.4; DB 4;
Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 29.4; DB 8;
Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                        Haemophilus influenzae complete genome sequence WO9633276-A1,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          40200210...
19-DEC-2002.
10-DEC-2002.
(UYLY-) UNIV LYON 1 BERNARD CLAUDE.
(BCOL-) ECOLE NORMALE SUPERIEURE DE LYON.
10.7%; Score 29.2; D
13.8%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.7%; Score 29.2; 3 52.5%; Pred. No. 87;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAS43104 standard; DNA; 325791 BP. Human Oestrogen receptor beta gene. WO200162793-A2.
                                                                                                                                                                                                                                                                                                         DNA; 1830121 BP
                                                                                                                                                                                                                                                     10.8%;
51.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10-MAY-2001.
(HUMA-) HUMAN GENOME SCI INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            19-JUN-2003.
(ISIS-) ISIS PHARM INC.
10-8%;
                                                                                         10.8%;
                                                                                                                                                                                                                                                                                                                                                                             (HUMA-) HUMAN GENOME SCI INC (UYJO ) UNIV JOHNS HOPKINS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Local Similarity 49.7%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        14-AUG-2003.
(GENE-) GENE LOGIC INC.
                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 590
ID ABQ83210 standard; DI
DE Human transnorts
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              30-AUG-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Local Similarity
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RESULT 595
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
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Best Local Similarity
                                                                          (PEKE ) PE CORP NY
                                                                                                                                                                                                                                                     Ouery Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                           AAT42063 standard;
                                                                                                                                                                                                                                                                                                                                                                 24-OCT-1996
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ID DE PN PD PA PA PA PA PA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABK14948 standard; DNA; 29793 BP.
Worm poly(ADP-ribose) glycohydrolase (PARG) sequence from cosmid F20C5.
US6337202-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABK14930 standard; DNA; 29793 BP.
Worm poly(ADP-ribose) glycohydrolase (PARG) sequence from cosmid F20C5.
US6333148-B1.
                                                                                                             ABLO4824 standard; cDNA; 6904 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 8954
WG200171042-A2.
27-SEP-2001.
                                                                                                                                                                                                                                                                                                                                                                                             ABL25924 standard; DNA; 7131 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 29245.
WO200171042-A2.
                                                                                                                                                                                                                                                                          84
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 29793;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   08-JAN-2002.
(KENT ) UNIV KENTUCKY RES FOUND.
ery Match 10.8%; Score 29.4; DB 6; Length 29793;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 29793;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 17073
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 29793
                                                         Length 6167;
                                                                                                                                                                                                       Length 6904;
                                                                                                                                                                                                                                                                                                                                               Length 7041;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               gene
                                                                                                                                                                                                                                                                          õ
                                                                                                                                                                                                                                                       ABN80067 standard; DNA; 7041 BP.

Human chemically modified disease associated gene SEQ ID 18 W0200200927-A2.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.8%; Score 29.4; DB 6; Length 7 st Local Similarity 47.1%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 8
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Poly adenosine diphosphate-ribose glycohydrolase (PARG)
US6395543-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL19434 standard; DNA; 17073 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID
W0200111042-A2.
27-SEP-2041.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABX14494 standard; DNA; 29793 BP.
Cosmid F20C5 containing C. elegans PARG genomic DNA.
19-SEP-2002.
(JACO/) JACOBSON M K.
(JACO/) JACOBSON E L.
(JAMEJ/) AME J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 29.4; DB 8;
Pred. No. 3.1e+02;
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(KENT ) UNIV KENTUCKY RES FOUND.

(KENT ) UNIV KENTUCKY RES FOUND.

(ery Match

-- " c'milarity 55.3%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.8%; Score 29.4; DB 6; 55.3%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.8%; Score 29.4; DB 4; 54.1%; Pred. No. 2.5e+02;
                                                         Score 29.4; DB 6;
Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.8%; Score 29.4; DB 4; 51.1%; Pred. No. 1.8e+02;
                                                                                                                                                                                                       10.8%; Score 29.4; DB 4; 51.1%; Pred. No. 1.8e+02;
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             WOZULL.
18-007-2001.
(EPIG-) EPIGENOMICS AG.
10.8%;
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Best Local Similarity
RESULT 585
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Best Local Similarity
RESULT 582
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Best Local Similarity
RESULT 584
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Best Local Similarity
RESULT 586
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RESULT 580
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                                                                        Best Local Similarity RESULT 579
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Length 110000;

Length 325791;

Length 245;

Length

Length 113000,

Length 110000,

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ADN02611 standard; DNA; 9227 BP.
S. dysenteriae serotype 8 O-antigen biosynthetic gene cluster (partial).
CN1429833-A.
                                                                                                                                                                                                                     Human polynucleotide associated with DNA replication SEQ ID NO 38. WO200177377-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABN80150 standard; DNA; 5937 BP.
Human chemically modified disease associated gene SEQ ID NO 167
WO200200927-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Chemically treated DNA repair gene fragment#64.

Chemically treated DNA repair gene fragment#64.

WO200181622-A2.

O1-NOV-2001.

A (EPIG-) EPIGENOMICS AG.

Query Match

10.7%; Score 29.2; DB 6; Length 73334;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 29.2; DB 2; Length 10607;
Pred. No. 2.5e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.7%; Score 29.2; DB 11; Length 9227; 53.5%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 5937;
                                                                                                                                                     Length 5501;
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           Length 5501
                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 5937;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 9110;
                                                         AAS61244 standard, DNA, 5501 BP.
Human gene regulation-associated gene oligonucleotide #199.
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABL34124 standard; DNA; 73334 BP.
Human immune system associated gene SEQ ID NO: 2097.
MU200200928-A2.
03-JAN-2002.
(RPIGT) EPIGENOMICS AG.
10.7%; Score 29.2; DB 6; Lenst Local Similarity 52.5%; Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   2401
       10.7%; Score 29.2; DB 6; 54.7%; Pred. No. 1.9e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 29.2; DB 6;
Pred. No. 2e+02;
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Pred. No. 1.9e+02;
                                                                                                                                                   Score 29.2; DB 6;
Pred. No. 1.9e+02;
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Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL34428 standard; DNA; 9110 BP,
Human immune system associated gene SEQ ID NO:
W0200200928-A2.
03-JAN-2002.
10-JAN-2002.
10.7%; Score 29.2; DB 6;
st Local Similarity 54.7%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 29.2; DB Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                        Human metastasis associated gene SEQ ID NO: WO200177376-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAV25594 standard; DNA; 10607 BP
                                                                                                                                                                                                      DNA; 5501 BP
                                                                                                                                                                                                                                                                                                                                              DNA; 5937 BP
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BOCT--2001.
(EPIG-) EPIGENOMICS AG.
GATO MATCh --: Larity 54.7%;
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.7%;
                                                                                               WOZULL.
18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
10.7%;
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 14-APR-1998.
(SYMB-) SYMBICOM AB.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       16-JUL-2003.
(UYNA-) UNIV NANKAI.
       Query Match
Best Local Similarity
RESULT 606
                                                                                                                                               Ouery Match
Best Local Similarity
RESULT 607
                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
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                                                                                                                                                                                                    ABL49338 standard;
                                                                                                                                                                                                                                                                                                                                           ABL34542 standard;
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Best Local Similarity
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RESULT 613
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL92255 standard; DNA; 5501 BP. Chemically treated DNA repair gene fragment complementary to#32.WO200181622-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 1161;
                                                                                                                                                                                                                                                                                         Length 1137;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 2123;
                                                                                                     05-FEB-2002.
(DOXU-) DOKURITSU GYOSEI HOJIN NOGYO SEIBUTSU SH.
(ery Match 10.7%; Score 29.2; DB 6; Length 895;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity 54.7%; Pred. No. 1.4e+02;
RESULT 602
ID AAD07821 standard; cDNA; 2529 BP.
B. Human secreted protein-encoding gene 13 cDNA clone HIBEB47, PP W0200132675-A1.
PD 10-MAY-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABNB1677 standard; DNA; 4220 BP.
Human CDK4 double bisulphite-converted genomic DNA SEQ ID NO
WC200236814-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 4220;
     DB 4; Length 820;
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Human CDK4 bisulphite-converted genomic DNA SEQ ID NO 1.
WO200236814-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                             Score 29.2; DB 10;
Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                      03-0CI-ZUUZ.
(BLIT-) ELITRA PHARM INC.
2ry Match 10.7%; Score 29.2; DB 8;
--- Tonal Similarity 65.2%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.7%; Score 29.2; DB 6; 53.5%; Pred, No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.7%; Score 29.2; DB 4; 62.2%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 29.2; DB 6;
Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 29.2; DB 6;
Pred. No. 1.8e+02;
                                                   ABK50746 standard; DNA; 895 BP.
Pyrus plant microsatellite DNA sequence #27.
JP2002034562-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABN60056 standard; DNA; 1767 BP.
Streptococcus polynucleotide SEQ ID NO 6025.
WO200234771-A2.
   10.7%; Score 29.2; 62.2%; Pred. No. 98;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABN96789 standard; DNA; 2123 BP.
Gene #3287 used to diagnose liver cancer.
WO200229103-A2.
                                                                                                                                                                                                                                                                                                                                   ADC91483 standard; DNA; 1161 BP.
E. faecium DNA sequence SEQ ID 1110.
US6583275-B1.
                                                                                                                                                                                             ACA33964 standard; DNA; 1137 BP. Prokaryotic essential gene #15621.03-021-2002.
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(GENO-) GENOME THERAPEUTICS CORP.
(ery_Match
.ery_match 10.7%; Scor.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WO20022.
11.APR-2002.
(GENE-) GENE LOGIC INC.
MATCh 10.7%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WCay-2002.
(EPIG-) EPIGENOMICS AG. 10.7%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CHIR-) CHIRON SPA.
(GENO-) INST GENOMIC RES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              01-NOV-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 597
ID ABK50746 standard, DN
DE Pyrus plant microsate
PN UP2002034582-A.
PD 05-FBB-2002.
PA (DOKU-) DOKURITSU GYC
                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 600
ID ARN69056 standard; DD DE Streptococcus polynuc
PN WC200234771-A2.
PA (CHIR-) CHIRON SPA.
PA (GENO-) INST GENOMIC
                                                                                                                                                        Best Local Similarity RESULT 598
                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 599
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RESULT 601
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                                                                                                                                           Query Match
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                                                                                                                                                                                                                                                                                                                                         PN DE
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ADO21878 standard; cDNA; 2655 BP.
Malaria parasite cyclic nucleotide phosphodiesterase PfPDE1 cDNA.
WO2004044192-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 1029;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 05-OCT-2000.
(HUWA.) HUWAN GENOME SCI INC.
(HUWA.) Match 10.6%; Score 29; DB 3; Length 1424;
lery Match 49.0%; Pred. No. 1.46+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 1205;
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                                                                                                                                                                                                                                        Length 694;
                                                      Length 694;
                                                                                                                                                                                                                                                                                                                                                                                                 10.6%; Score 29; DB 8; Length 855; 54.1%; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABZ16971 standard; DNA; 1573 BP.
Arabidopsis thaliana stress regulated gene SEQ ID NO 4776.
WO200216655-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACF70507 standard; DNA; 1029 BP.
Photorhabdus luminescens nucleotide sequence #8974.
WO200294867-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human ovarian antigen HE6CR19 CDNA, SEQ ID NO:721
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WO200110...
28-FEB-2002.
(SCRI ) SCRIPPS RES INST.
(SYGN ) SYNGENTA PARTICIPATIONS AG.
(SYGN ) SYNGENTA PARTICIPATIONS AG.

"" Match """ "" " 48.0%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADM19639 standard; cDNA; 1502 BP.
Novel human channel/transporter gene #192 clone
WO200154472-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WOACCE...
MOACCE...
(30-JAN-2002.
(HUMAL) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 10.6%; Score 29; DB 5; I
                                           vuery Match 10.6%; Score 29; DB 2; 1
Best Local Similarity 58.8%; Pred. No. 1.1e+02;
RESULT 625
                                                                                                            AAX80476 standard; cDNA; 694 BP.
Mouse INK-4 protein p18 encoding cDNA.
US591997-A.
(COLD-) COLD SPRING HARBOR LAB.
(YESH) UNIV YESHIVA BINGTEIN COLLEGE.
YESH) WALCH
58.88; Pred. No. 1.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.6%; Score 29; DB 10; 54.1%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAC74396 standard; cDNA; 1424 BP.
Human secreted protein gene 1 SEQ ID NO:11.
WO200058496-A1.
D 15-AUG-1996.
A (SJUD-) ST JUDE CHILDREN'S RES HOSPITAL.
Query Match
                                                                                                                                                                                                                                                                                                                                                             03-JAN-2003.
(SYGN ) SYNGENTA PARTICIPATIONS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           RESULT 633
ID ABX09864 Btandard; DNA; 2814 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      28-NOV-2002.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                                                                                                                 ADA72899 standard; DNA; 855 BP.
Rice gene, SEQ ID 6225.
WO2003000898-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  27-MAY-2004.
(TANA ) TANABE SEIYAKU CO.
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Best Local Similarity
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADEILI69 standard; DNA; 394191 BP.
Human transporter protein encoding gene SEQ ID NO:1.
Ery Match
St. Local Similarity 54.7%; Pred. No. 5.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 95400
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 110000
                                                                                                                                                                                                                                                                                                                                                                                                                          Length 75899
                                                                                                                                                                                                                                                             Score 29.2; DB 6; Length 75899; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AUI13990 standard; DNA; 75899 BP.
Human protein phosphatase 1B (PTP1B) genomic DNA SeqID 243
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.6%; Score 29; DB 6; Length 257; 58.8%; Pred. No. 74;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               4; Length 408;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.6%; Score 29; DB 8; Length 553; 57.0%; Pred. No. 98;
                                   3SULT 615

ABK85261 standard; DNA; 75899 BP.

A BAR85261 standard; DNA for protein phosphatase 1B, PTP1B.

US2002055479-A1.

0 09-MAY-2002.

1 (COWA); COWSERT L M.

4 (WYAT/) WYATT J.

A (FREI/) FREIER S M.

A (MOXIA) MOXIA B P.

A (BUIL/) BUILER M M.

A (BUIL/) BUILER M M.

A (MOXIA) MCKAY R.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABN26336 standard; CDNA; 257 BP.
Human ORFX polynucleotide sequence SEQ ID NO:21149.
WO200192523-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                       DE Human laminin alpha 4 (LAWA4) genomic DNA.

Query Match
10.7%; Score 29.2; DB 12;

Query Match
54.7%; Pred. No. 5.40+02;
                                                                                                                                                                                                                                                                                                                                                                          US.C.C.-2003.
27-MOV-2003.
(ISIS-) ISIS PHARM INC.
(EXY Match 10.7%; Score 29.2; DB 12;
(EXY Match 10.7%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Aspergillus oryzae polymucleotide SEQ ID NO 3312 WO200279476-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.7%; Score 29.2; DB 6; 53.5%; Pred. No. 5.7e+02;
                    52.5%; Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY
(NARE-) NAT RES INST BREWING.
(NORQ ) NAT FOOD RES INST MIN AGRIC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB /
88;
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Human prostate expression marker cDNA 3814.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 29;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human polynucleotide SEQ ID NO 995. W0200164835-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAT36042 standard; cDNA; 694 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABZ54199 standard; cDNA; 553 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.6%;
51.1%;
                                                                                                                                                                                                                                                             10.7%;
50.7%;
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Query Match

Best Local Similarity

RESULT 623

ID ABZ$4199
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               06-DEC-2001.
(CURA-) CURAGEN CORP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 624
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Best Local Similarity
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Best Local Similarity
RESULT 622
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Best Local Similarity
RESULT 619
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Best Local Similarity
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RESULT 616
                       Best Local Similarity
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ABL70514 standard; DNA; 15674 BP. Chemically treated cell signalling DNA sequence complementary to#202.
WO200202807-A2.
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(ADSC.) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.

10.6%; Score 29; DB 10; Length 349881;
tr Match
T7.8%; Pred. No. 9.7e+02;
                                                                                      Score 29; DB 6; Length 15674; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (HUMA-) HUMAN GENOME SCI INC.

ery Match 10.5%; Score 28.8; DB 4; Length 706;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABV94448 standard; cDNA; 396 BP.
Breast carcinoma related nucleotide sequence SEQ ID NO:439-
W0200246467-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAK87618 standard; DNA; 706 BP.
Human immune/haematopoietic antigen genomic sequence SEQ
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABK76543 standard; DNA; 318 BP.
Bacillus licheniformis genomic sequence tag (GST) #3834-WO200229113-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Osteoarthritis-associated polymorphic nucleotide #181 W02003054166-A2.
03-JUL-2003.
(INCY-) INCYTE GENOMICS INC.
ery Match
st Local Similarity 61.0%; Pred. No. 6.9e+02;
                                                                                                                                                                                                                                                                                                Photorhabdus luminescens nucleotide sequence #27
WO200294867-A2.
                                                                                                                                                                                                                                                                                                                  DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      9
 ABL34477 standard; DNA; 15674 BP.
Human metastasis associated gene SEQ ID NO: 30.
W0200177376-A2.
W0200177376-A2.
(EPIGC-2001.
(EPIG-) EPIGENOMICS AG.
10.6%; Score 29; DB 6; St Local Similarity 55.4%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                             10.6%; Score 29; DB 6; 55.4%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 29; DB 6;
Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 10;
6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.5%; Score 28.8; DB 61.8%; Pred. No. 1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.5%; Score 28.8; 1 50.7%; Pred. No. 93;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 29;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADC86642 standard; DNA; 349881 BP.
Human GPCR gene SEQ ID NO:1095.
EP1270724-A2.
02-JAN-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DNA; 127098 BP
                                                                                                                                                                                                                                                                                  ACF65374 standard; DNA; 69727 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (NOVO ) NOVOZYMES BIOTECH INC. (NOVO ) NOVOZYMES AS.
                                                                                                                                                                                                                                                                                                                                                                                                                                         10.6%;
63.8%;
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54.1%;
                                                                                                                                                                                             10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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Best Local Similarity
RESULT 647
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Best Local Similarity
RESULT 651
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(IPSO-) IPSOGEN.
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                                                                                            Query Match
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                                                                                                                                                                                                                                                                                           AAD05229 standard; cDNA; 3143 BP.
Human secreted protein-encoding gene 10 cDNA clone HHEPJ23, SEQ ID NO:20.
WO200134629-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Drosophila melanogaster expressed polynucleotide SEQ ID NO 17426 WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 29; DB 6; Length 15674; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 3572;
N. lactamica DNA encoding a vaccine antigen #16.
W0200277648-A2.
03-OCT-2002.
(MICR-) MICROBIOLOGICAL RES AUTHORITY.
(IMCO-) IMPERIAL COLLEGE INNOVATIONS LTD.
ery Match
st Local Similarity 52.0%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                              Length 3125
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                                                                                                                                                                                                                                                                                                                                          TY-MAY-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 10.6%; Score 29; DB 4; Length 3143;
ery Match 50.4%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 5046;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.6%; Score 29; DB 5; Length 5046; 52.0%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.6%; Score 29; DB 4; Length 5970; 47.1%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                A. thaliana APETALA2 (AP2) complete genomic sequence W09941974-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                ADF59725 standard; cDNA; 3572 BP.
Human contig polynuclectide sequence SEQ ID NO:2092
WO2003080795-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL32363 standard; DNA; 15674 BP.
Human immune system associated gene SEQ ID NO: 336
WO200200928-A2.
                                                                                                                                                                                                                                              10.6%; Score 29; DB 10; 50.4%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.6%; Score 29; DB 10; 52.0%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.6%; Score 29; DB 5; 52.0%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 29; DB 2;
Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADF67944 standard; DNA; 5046 BP.
T. thermophila triterpenoid cyclase DNA #2.
DE19957889-A1.
                                                                                                                              Human cDNA encoding clone SPLEN20136700.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAH48716 standard; DNA; 5046 BP.
T. thermophila triterpenoid cyclase DNA
EP1130103-A2.
                                                                                                                                                                                      07-MAY-2003.
(HELL-) HELIX RES INST.
(REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA: 11721 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL07648 standard; cDNA; 5970 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (CELA ) CELANESE VENTURES GMBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WO99412, 26-AUG-1999. (REGC ) UNIV CALIFORNIA. 10.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.6%;
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                                                                                                     Best Local Similarity RESULT 634
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RESULT 642
                                                                                                                                                                                                                                                            Best Local Similarity RESULT 635
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Best Local Similarity
RESULT 639
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(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 640
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Best Local Similarity
RESULT 638
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(AXIV-) AXIVA GMBH.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAZ07666 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        02-OCT-2003 (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                05-SEP-2001
                                                                                          Query Match
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Length 127098;

Length 110000,

Length 69727;

Length 15674;

Length 110000;

NO:42430.

ΠD

Length 318;

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AAS45499 standard; DNA; 5476 BP.
Chemically pretreated complementary DNA associated with cell cycle #102.
W0200168911-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL49381 standard; DNA; 5476 BP.
Human polynucleotide associated with DNA replication SEQ ID NO 81.
NO 20017777-A2.
                                                    AAZ49290 standard; DNA; 2849 BP.
Homology vector 443-88.14 2.8 kb FPV genomic sequence, SEQ ID
US6001369-A.
                                                                                                                                                                                                                                                                                                                                  10.5%; Score 28.8; DB 3; Length 2849; 48.8%; Pred. No. 2e+02;
                                                                                                                                                  10.5%; Score 28.8; DB 3; Length 2849; 48.8%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADQ25278 standard; DNA; 5127 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 8098
WQ2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABM80238 standard; DNA; 15121 BP.
Human chemically modified disease associated gene SEQ
WO200200927-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL34114 standard; DNA; 5476 BP.
Human immune system associated gene SEQ ID NO: 2087
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human immune system associated gene SEQ ID NO: 1722.
                                                                                                                                                                                                                                                                                                                                                                                            AAS87447 standard; cDNA; 2856 BP.
DNA encoding novel human diagnostic protein #23251.
WO200175067-A2.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
PRY MATCH 10.5%; Score 28.8; DB 12;
it Local Similarity 48.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 28.8; DB 5;
Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 28.8; DB 4;
Pred. No. 2.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 28.8; DB 6;
Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 28.8; DB 6;
Pred. No. 2.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 28.8; DB 6;
Pred. No. 2.6e+02;
Best Local Similarity 48.8%; Pred. No. 2e+02; RESULT 661
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            MOCOULD
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.5%; Score 28.8;
                                                                                                                                                                                                AACG7857 standard; DNA; 2849 BP.
FPV homology vector 749-75.82 fragment 1.
US6136318-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          WO2001.
18-007-2001.
(EPIG-) EPIGENOMICS AG.
10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.5%;
56.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WGSCG-2002.
(BPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         03-JAN-2002.
(BIG-) EPIGENOMICS AG.
Query Match 10.5%;
Best Local Similarity 50.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          53.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                            14-DEC-1999.
(SYTR ) SYNTRO CORP.
                                                                                                                                                                                                                                                                           24-OCT-2000.
(JUNK/) JUNKER D E.
(COCH/) COCHRAN M D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 667
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
                                                                                                                                                                       Local Similarity
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                                                                                                                                                        Query Match
                                                                                                                                                                                                                                                                                                                                                         Best Loca
RESULT 663
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          RESULT 668
                                                                                                                                                                             Best Loc
RESULT 662
                  AAK87616 standard; DNA; 717 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:42428.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match 10.5%; Score 28.8; DB 2; Length 2849;
Best Local Similarity 48.8%; Pred. No. 2e+02;
RESULT 658
DE AAQ70573 standard; DNA; 2849 BP.
DE Homology vector 443-88.14 contg. fowlpox virus genomic fragment.
PN W09419015-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 RESULT 656

ID ABL28530 standard; DNA; 2414 BP.

DD Drosophila melanogaster genomic polynucleotide SEQ ID NO 37063.

PD 75EP-2001.

PA (PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AQG8937 standard; DNA; 2849 BP.

Fowlpox virus (FPV) genomic EcoRI fragment on vector 443-88.14.

W09419014-A1.

01-SEP-1994.

(SYTR ) SYNTRO CORP.

(GARG) NIPPON ZEON KK.

10.5%; Score 28.8; DB 2; Length 2849;

EL Docal Similarity 48.8%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                        ABK82130 standard; cDNA; 1095 BP.

DNA encoding novel floral meristem identity protein LpMADS4-2
WO200233091-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 2849;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 2414;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 2849;
                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 1095;
                                                                                                                     717;
                                                                                                                                                                                                                                                                               Length 993;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Homology vector 443-88.14 containing FPV genomic fragment.
USS925358-A.
D 20-JUL-1999.
A (SYTR ) SYNTRO CORP.
10.5%; Score 28.8; DB 2: Lienath 28.
                                                                                                                   Length
                                                                                                                                                                           ABQ75352 standard; cDNA; 993 BP.
Human lung specific nucleic acid sequence SEQ ID NO:91
WO200264788-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 28.8; DB 10;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.5%; Score 28.8; DB 4; 50.7%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.5%; Score 28.8; DB 2; 48.8%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.5%; Score 28.8; DB 6; 48.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                             10.5%; Score 28.8; DB 6; 54.8%; Pred. No. 1.4e+02;
                                                                                                                   10.5%; Score 28.8; DB 4; 51.6%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Enterococcus faecalis polynucleotide #1947.
US6617156-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                     (AGRI-) AGRIC VICTORIA SERVICES PTY LTD (AGRE-) AGRESEARCH LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAT48512 standard; DNA; 2849 BP. Homology vector 443-88.14.
19-086-41.
(SYTR ) SYNTRO CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAX81142 standard; DNA; 2849 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADH84062 standard; DNA; 1296 BP
                                                                                                (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (DOUC/) DOUCETTE-STAMM L A. (BUSH/) BUSH D.
                                                                                                                                                                                                                                          22-AUG-2002.
(DIAD-) DIADEXUS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            01-SEP-1994.
(SYTR ) SYNTRO CORP.
                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 657
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                                                                                                                                 Best Local Similarity RESULT 653
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      09-SEP-2003
                                                                                                                                                                                                                                                                                                                                                                                                     25-APR-2002
                                                                               09-AUG-200
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                                                                                                                     Query Match
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Length 5127

Length 2856;

PN PA PA

NO 255

ΩI

Length 11805

Length 5476;

Length 5476;

Length 5476

DB 6; Length 15121;

Length 177851;

Length 349980;

Length 426;

Length 450;

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ABK62981 standard; cDNA; 426 BP.
Rat sequence differentially expressed in response to a hepatotoxin #888.
WO200210453-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABN92575 standard; DNA; 450 BP.
Staphylococcus epidermidis ORF nucleic acid sequence SEQ ID NO:2038.
US6380370-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABS64610 standard; cDNA; 454 BP.
Human cDNA probe selectively hybridising to foetal cell mRNA #25.
W0200255985-A2.
       10.5%; Score 28.8; DB 11; Length 128361; 56.2%; Pred. No. 8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human colon cancer related nucleotide sequence SEQ ID NO:484.
WO200229086-A2.
II.ARR-2002.
Y Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 28.6; DB 6; Length 651; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABN63352 standard; cDNA; 518 BP.

Human cancer related polynucleotide SEQ ID NO 3319.

W2020214500-A2.
21-FEB-2002.

(CHIR ) CHIRON CORP.

(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                             (CNRS ) CNRS CENT NAT RECH SCI.
(IFRE-) IFREMER INST FR RECH EXPL MER.
10.5%; Score 28.8; DB 5;
t Local Similarity 54.8%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.5%; Score 28.6; DB 6; 49.0%; Pred. No. 1.2e+02;
                                                                                                                                              8
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(HOFF) ROCHE DIAGNOSTICS CORP.
ery Match 10.5%; Score 28.6; DB 6;
ery Match 49.0%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                           PA (GENE-) GENE LOGIC INC.
Query Match
Best Local Similarity 67.8%; Pred. No. 1.2e+02;
RESULT 682
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.5%; Score 28.6; DB 6; 51.1%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.5%; Score 28.6; DB 6; 47.5%; Pred. No. 1.4e+02;
                                                             AAL57272 standard, DNA; 177851 BP.
bA438B23-1 'human modifier of p53 pathway' DNA
                                                                                                                                                                                                                #2
                                                                                                                                                                                                              Pyrococcus abyssi genomic fragment FR2792651-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           30-APR-2002.
(GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABQ98890 standard; DNA; 651 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human ORF697 coding sequence.
US2002082206-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.5%;
51.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  27-JUN-2002.
(LEAC/) LEACH M D.
(MEHR/) MEHRABAN F.
(CONL/) CONLEY P B.
(TOPP/) TOPPER J N.
(LAWD/) LAW D.
 Query Match
Best Local Similarity
RESULT 679
                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity
RESULT 684
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 686
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 685
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                                                                                                                                                                                                                                                                                                                                                                                                        07-FEB-2002
                                                                                                                                                                                                                                                 27-OCT-20(
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                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAK71622 standard; DNA; 27062 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:26434.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                28.8; DB 12; Length 112604; No. 7.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 02-JAN-2003.

(NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.

(ADSC.) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.

ETY Match 10.5%; Score 28.8; DB 10; Length 29221;

cery Match 48.8%; Pred. No. 4.7e+02;
                                                                                                                                                                                            86
                                                                                                                            Length 16602;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human testicular antigen encoding DNA fragment SEQ ID NO: 2867 WO200155317-A2.
                                                                                                                                                                                                                                                                                                                                                         13-JUN-2002.
(EPIG-) EPIGENOMICS AG.
(ery Match 10.5%; Score 28.8; DB 6; Length 18357;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 27062;
                                                                                                                                                                                                                                                             Length 16602
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 27062;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAL05346 standard; DNA; 27062 BP.
Human reproductive system related antigen DNA SEQ ID NO: 8034.
WO200155320-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 4; Length 27062;
                                                                                                                                                 .....vvvvy standard; DNA; 16602 BP.

Human chemically modified disease associated gene SEQ ID NO W0200200927-A2.
                                                                                                                                                                                                                                                                                                          ABQ67084 standard; DNA; 18357 BP.
Human angiogenesis associated polynucleotide SEQ ID NO 114.
WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADQ18153 standard; DNA; 112604 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 970.
WO2004048938-A2.
                              ABL32727 standard; DNA; 16602 BP.
Human immune system associated gene SEQ ID NO: 700.
W0200200928-A2.
03-JAN-2002.
(RPIG-) EPIGENOMICS AG.
10.5%; Score 28.8; DB 6; Le
st Local Similarity 52.5%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                      (EPIG-) EPIGENOMICS AG.
10.5%; Score 28.8; DB 6;
it Local Similarity 52.5%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.5%; Score 28.8; DB 4; 54.8%; Pred. No. 4.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.5%; Score 28.8; DB 4; 54.8%; Pred. No. 4.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   02-401.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
10.5%; Score 28.8; DB 4;
ery Match
54.8%; Pred. No. 4.6e+02;
     No. 3.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADN95229 standard; DNA; 128361 BP.

Human BEC/LEC-related gene sequence SeqID151.
WO2003080640-A1.
02-OCT.
02-OCT.
UDDW-) LUDWIG INST CANCER RES.
(LICH) LICENTIA LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.5%; Score 62.5%; Pred.
   56.2%; Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADC86182 standard; DNA; 29221 BP.
Human GPCR gene SEQ ID NO:635.
BP1270724-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABL98215 standard; DNA; 27062 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 673
ID AAK11622 standard; DN
DE Human immune/haematop
PN W0200157182-A2.
PD 09-AUG-2001.
PA (HUMA-) HUMAN GENOME
                                                                                                                                     Best Local Similarity RESULT 671
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 676
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 677
Best Local Similarity RESULT 670
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                          Best Local Similarity
RESULT 672
                                                                                                                                                                                                                                                           Query Match
                                                                                                                        Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           RESULT 675
ID ABL982
DE Human PN WO2001:
PD 02-AUG-PA (HUMA-)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               RESULT 678
                                                                                                                                                                                                                                                                                                              DE DE LO
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Length 454;

Length 648;

Length 518;

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Length 1552;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 2000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.5%; Score 28.6; DB 4; Length 1813; 55.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                    Length 1650;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 1819;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 2037;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 4; Length 2243;
                                                                                                                                                                                                                                                                                                                                                                                          Length 1765;
                ADQ22201 standard; DNA; 1552 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 5021.
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 28.6; DB 12;
Pred. No. 2.1e+02;
                                                                      10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
PLY MATCH 10.5%; Score 28.6; DB 12;
st Local Similarity 52.6%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 28.6; DB 8;
Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                            AAH99127 standard; cDNA; 1813 BP.
Human EST-derived coding sequence SEQ ID NO: 984
WO200154477-A2.
                                                                                                                                                                                                                                                                                                                                                                                            8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAH29911 standard; DNA; 2243 BP.
C albicans apoptosis associated coding sequence
WO200102550-A2.
                                                                                                                                                                                                                                                  10.5%; Score 28.6; DB 8; 59.0%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 28.6; DB 4;
Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                            Score 28.6; DB
Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.5%; Score 28.6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human cDNA sequence SEQ ID NO:14479.
EPI074617-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human novel polynucleotide #446. WO200274961-Al.
                                                                                                                                                                                                                                                                                            Human novel polynucleotide #447.
26-SPB.-Act.
                                                                                                                                              Prokaryotic essential gene #3470.
W220271183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADJ40941 standard; cDNA; 2000 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.5%;
51.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.5%;
51.1%;
                                                                                                                                                                                                                                                                                                                                                                                            10.5%;
51.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PD 07-FEB-2001.

PA (HELL-) HELIX RES INST.

QUETY MATCh 10.5%;

Best Local Similarity 51.1%;

RESULT 703
                                                                                                                                                                                                               03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     11-JAN-2001.
(JANC ) JANSSEN PHARM NV.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       BRIGGS S P.
COOPER B.
GLAZEBROOK J.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (BRIG/) BRIGGS S P.
(COOP/) COOPER B.
(GLAZ/) GLAZEBROOK J.
(GOFF/) GOFF S A.
(KATA/) KATAGIRI F.
(KREP/) KREPS J.
(FROY/) PROVART N.
(RICK/) RICKE D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     22-JAN-2004.
(BUDW/) BUDWORTH P.
(MOUG/) MOUGHAMER T.
                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 699
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Local Similarity
                                                                                                                                                                                                                                                                                                                                                       26-SEP-2002.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  02-AUG-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Plant cDNA #1941.
US2004016025-A1.
                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
       RESULT 696
                                                                                                                                                                                                                                                                                       RESULT 698
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         RESULT 702
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAA01909 standard; cDNA; 831 BP.
Human colon cancer cell line polynucleotide sequence SEQ ID NO:1900.
WO9958675-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 1367
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               21-SEP-2000.
21-SEP-2000.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
10-5%; Score 28.6; DB 3; Length 1509;
10-7, Similarity 51.1%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.5%; Score 28.6; DB 3; Length 1483; 51.1%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     antigen nucleotide sequence SEQ ID NO:159
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 855;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.5%; Score 28.6; DB 4; Length 859; 51.1%; Pred. No. 1.5e+02;
                                                                                        Length 736;
                                                                                                                                                                                                                                   Length 773;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 855,
                                                                                                                                                                                                                                                                                                                                                                                            Length 831
                                                                                                                                                                                                                                                                                                                                                                                                                                              ABZ16462 standard; DNA; 855 BP.
Arabidopsis thaliana stress regulated gene SEQ ID NO 4267.
WO200216655-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAC77532 standard; cDNA; 1483 BP.
Human ORFX ORF3087 polynucleotide sequence SEQ ID NO:6173
WO200058473-A2.
AAH34681 standard; cDNA; 736 BP.
Human colon cancer antigen encoding cDNA SEQ ID NO:1763
WO200122920-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human soft tissue sarcoma-upregulated DNA - SEQ ID 5457
WO2004048938-A2.
                                                                                                                                        AAL25096 standard; cDNA; 773 BP.
Human breast cancer expressed polynucleotide 17553
WO200151628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.5%; Score 28.6; DB 12; 52.0%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     WOZUWALUZA

28-FEB-2002.

SCRI ) SCRIPPS RES INST.

(SYGN ) SYNGENTA PARTICIPATIONS AG.

(SYGN ) SYNGENTA PARTICIPATIONS AG.

10.5%; Score 28.6; DB 6;
                                              MCZVOLL.

05-APR-2001.

(HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

10.5%; Score 28.6; DB 4;

ery Match

54.5%; Pred. No. 1.4e+02;
                                                                                                                                                                                            19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.5%; Score 28.6; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (SYGN ) SYNGENTA PARTICIPATIONS AG.

ry Match
10.5%; Score 28.6; DB 8;
t Local Similarity 57.1%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                            Score 28.6; DB 3;
Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAHOS511 standard; cDNA, 859 BP.

Human cDNA clone (5'-primer) SEQ ID NO:2346.

BP10/4617-A2.

(77-FEB-2001.

(HELI-) HELIX RES INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADA69101 standard; DNA; 855 BP.
Arabidopsis thaliana gene, SEQ ID 2424.
WO2003000898-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    CDNA; 1509 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADQ22637 standard; DNA; 1367 BP
                                                                                                                                                                                                                                                                                                                                                                                            10.5%;
53.9%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           05-OCT-2000.
(CURA-) CURAGEN CORP.
                                                                                                                                                                                                                                                                                                                                                         (CHIR ) CHIRON CORP. (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 692
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Human colon cancer
WO200055351-A1.
                                                                                                                                                                                                                               Query Match
Best Local Similarity
RESULT 689
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Ouery Match
Best Local Similarity
SULT 693
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 694
                                                                                          Query Match
Best Local Similarity
RESULT 688
                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 690
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 691
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Best Local Similarity
RESULT 695
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAC98149 standard;
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                         18-NOV-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               03-JAN-2003
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Length 4711;

Length 4750;

Length 5236;

PN PR

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Drosophila melanogaster expressed polynucleotide SEQ ID NO 40028 WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human chemically modified disease associated gene SEQ ID NO 225-W0200200927-A2.
                                                                                                                                                                           Score 28.6; DB 10; Length 4711; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human MNB modulator of the SREBP pathway DYRKIA gene sequence. WO2003066811-A2.

14.AUG-2003.

(EXEL-) EXELIXIS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADA20363 standard; DNA; 6246 BP.
Prostate tumour related genomic DNA complement sample #14 WO2002103042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABX33966 standard; DNA; 6246 BP.
ABX33966 standard; DNA; 6246 BP.
WO200202808-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AUA84170 standard; DNA; 6246 BP.
Human renal/prostate carcinoma associated DNA SEQ ID
WO2002103041-A2.
                     Score 28.6; DB 10;
Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28.6; DB 10;
Pred. No. 3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 28.6; DB 6;
Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                10.5%; Score 28.6; DB 4; 53.0%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 28.6; DB 6;
Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 28.6; DB 6;
Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28.6; DB 8;
Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 28.6; DB 8;
Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                               ABL32350 standard; DNA; 5236 BP.
Human immune system associated gene SEQ ID NO:
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                     ADE62659 standard; DNA; 4711 BP.
Human gene XM 042066, SEQ ID NO 8591
WO2003016475-Ã2.
                                                                                                                                                                                                                              ABL15182 standard; cDNA; 4750 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABL33017 standard; DNA; 6246 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DNA; 7323 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Wodoucou
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       WOZOWZ-
10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Wolvostra
27-DEC-2002.
(EPIG-) EPIGENOMICS AG.
10.5%;
                       10.5%;
                                                                                                                                                                           10.5%;
54.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.5%;
51.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.5%;
57.1%;
                                                                                                                                      (GEHO ) GEN HOSPITAL CORP. (FARB ) BAYER AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      27-DEC-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                 03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                              27-SEP-2001.
(PEKE) PE CORP NY.
                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 715
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 720
                     Query Match
Best Local Similarity
RESULT 713
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABN80208 standard;
                                                                                                                                                                                          Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
       (FARB ) BAYER AG.
                                                                                                                        27-FEB-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Ouery Match
                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
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                                                                                                                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          RESULT 716
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAD12567 standard; cDNA; 3323 BP.
Human protein having hydrophobic domain encoding cDNA clone HP03700.
WC200149728-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABS64611 standard; cDNA; 3661 BP. Human cDNA probe selectively hybridising to foetal cell mRNA #26 W0200255985-A2. 18-JUL-2002.
                                                                                                                           Length 2735;
                                                                                                                                                                                             685.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAH54843 standard; DNA; 3054 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:4207
WO200134809-A2.
(GLAX ) GLAXO GROUP LID.
                                                                                                                                                                                                                                                                Length 2743
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 3054;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAH54548 standard; DNA; 3126 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:3912
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 4263;
                                                                                                                                                                                                                                                                                                                                                                                                                      Length 3013;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 3661;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 3126;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 3323
                                                                                                                                                           AUF77010 standard; cDNA; 2743 BP.
Novel human secreted and transmembrane protein cDNA SeqID
                                                                                                                                                                                                                                                                                                           AAF18334 standard; DNA; 3013 BP.
Lung cancer associated polynucleotide sequence SEQ ID 353
WO200055180-A2.
                                                                                                                         10.5%; Score 28.6; DB 12; 49.0%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                      04-5EP-2003.
(GETH ) GENENTECH INC.
(ery Match 10.5%; Score 28.6; DB 10;
(ery Match 49.0%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 28.6; DB 3;
Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         LTD.
10.5%; Score 28.6; DB 4;
49.0%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.5%; Score 28.6; DB 4; 49.0%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 28.6; DB 6;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.5%; Score 28.6; DB 6; 49.0%; Pred. No. 2.6e+02;
       50.4%; Pred. No. 2.2e+02;
                                   ADL12643 standard; cDNA; 2735 BP.
Human steroid-induced C3A liver cell cDNA #372.
US6673549-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABZ32319 standard; DNA; 4263 BP.
Candida albicans essential gene SEQ ID NO 6606.
WO200253728-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADE62655 standard; DNA; 4711 BP.
Human gene XM 042066, SEQ ID NO 8587.
W02003016475-Ā2.
27-FEB-2003.
(GEHO ) GEN HOSPITAL CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (HOFF ) ROCHE DIAGNOSTICS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.5%;
                                                                                                                                                                                                                                                                                                                                                                                (HUMA-) HUMAN GENOME SCI INC (ROSE/) ROSEN C A.
                                                                                                                                                                                                                                                                                                                                                                                                                   10.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
                                                                                                                                                                                            Bet 2013-A2.

2013-A2.

2014-A GENENTECH 1.

2017 MAtch
Best Local Similarity 4
RESULT 706
ID AAF18334 standar?
ID AAF18334 standar?
PB Lung cancer
PW WO20005517
PD 21-SEP
PA (HT
Best Local Similarity
RESULT 704
ID ADLISCA standard; cl
DE Human steroid-inducee
PN US6673549-B1.
PD 06-JAN-2004
PA (INCY-) INCYTE CORP.
                                                                                     06-JAN-2004.
(INCY-) INCYTE CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity
RESULT 710
                                                                                                                                  Best Local Similarity RESULT 705
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Best Local Similarity
RESULT 712
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 711
                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                           Query Match
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PNE

PNE

PAPE

ID DE PN PD PD

Length 6246;

990

Length 6246;

Length 6246;

Length 6246;

Length 35586;

Length 38045;

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AAKB2012 standard, DNA, 49561 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36824
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAL23194 standard; cDNA; 371 BP.
Human breast cancer expressed polynucleotide 15651
WO200151628-A2.
                                                  Score 28.6; DB 12;
Pred. No. 5.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.6; DB 10;
Pred. No. 8.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 28.6; DB 10;
Pred. No. 8.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 28.6; DB 10;
Pred. No. 8.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 28.6; DB 10;
No. 8.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 28.6; DB 9;
Pred. No. 6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 28.6; DB 6;
Pred. No. 8.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.4%; Score 28.4; DB 5; 54.9%; Pred. No. 1.2e+02;
                                                                                                                                                                                                                                                                                                                               10.5%; Score 28.6; DB 4; 51.1%; Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                  10.5%; Score 28.6; DB 2; 48.0%; Pred. No. 7.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                     10.5%; Score 28.6; DB 2; 57.1%; Pred. No. 8.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human prostate expression marker cDNA 56456.
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABA92787 standard; DNA; 640681 BP.
Buchnera sp. genomic DNA SEQ ID NO:1.
J92001292771-A.
23-OCT-2001.
IKRA) RIKAGAKU KENKYUSHO.
ery Match
Local Similarity 54.2%; Pred. No
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Pred.
                                                                                                   AAD56159 standard; DNA; 38045 BP.
Human salvador genomic DNA.
WO2003035845-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CDNA; 261 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ACH49514 standard; cDNA; 416 BP.
Human leukocyte cDNA #1108.
US2003073623-A1.
                                                                                                                                                                                                                                                                                             09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                ULMAY-2003.
(GEHO ) GEN HOSPITAL CORP.
(ery Match (ery Match (ery Match))
                                                  10.5%;
54.2%;
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57.1%;
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57.1%;
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55.6%;
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57.1%;
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STACHE-CRAIN B.
DICKSON M C.
JONES L W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             17-APR-2003.
(DRMA/) DRMANAC R T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
                                                                                                                                                                                                     Best Local Similarity
RESULT 732
EP1426442-A1.
09-JUN-2004.
(MTMM-) MTM LAB AG.
                                                               Best Local Similarity RESULT 731
                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
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Best Local Similarity
RESULT 734
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Best Local Similarity
RESULT 735
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Best Local Similarity
RESULT 736
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Best Local Similarity
RESULT 737
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABV56465 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
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Best Local Similarity
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(STAC/)
(DICK/)
(JONE/)
                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
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DE PR
PR
PR
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Human dentin sialophosphoprotein precursor (DSPP) gene SEQ ID NO:1.
WO200258722-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Genomic sequence #314 encoding human colon associated polypeptide WO200155302-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAK90312 standard; DNA; 10766 BP.
Human digestive system antigen genomic sequence SEQ ID NO: 3888.
WO200155314-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.5%; Score 28.6; DB 9; Length 10766; 51.1%; Pred. No. 3.8e+02;
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(HUMA-) HUMAN GENOME SCI INC.
ery Match
ery Match
'...a' Similarity 51.1%; Pred. No. 3.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 10766
                                                                                                                                                                                                                                                                                                                   Length 7746;
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                     Length 7323;
                                                                                                                                                                           Length 7560;
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Human novel colon related polypeptide DNA SEQ ID NO 792.
US2003050231-A1.
                                                                                                                                                                                                                            ABK40048 standard; DNA; 7746 BP.
Human chemically pretreated gene sequence #65 strand
W0200202806-A2.
10-JAN-2002.
(EPIGS) EPIGENOMICS AG.
10-St Actor Section 10.5%; Score 28.6; DB 6; Leng
st Local Similarity 48.0%; Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                  ABL33857 standard; DNA; 7746 BP.
Human immune system associated gene SEQ ID NO: 1830.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                01-AGG-2002.
(SHAN-) SHANGHAI RES CENT BIOTECHNOLOGY.
(ery Match 10.5%; Score 28.6; DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      WC.Z.C.C.
02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 10:5%; Score 28.6; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PA (USSH ) US DEPT HEALTH & HUMAN SERVICES.
Query Match 10.5%; Score 28.6; DB 3;
Best Local Similarity 51.1%; Pred. No. 3.5e+02;
RESULT 726
                                                                                                                                                                           10.5%; Score 28.6; DB 6; 54.2%; Pred. No. 3.4e+02;
                   10.5%; Score 28.6; DB 6; 45.7%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28.6; DB 6;
Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAA88864 standard; DNA; 8201 BP.
Human dentin sialophosphoprotein gene.
WO200062065-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADO34038 standard; DNA; 35586 BP.
Human CLA2 gene, SEQ ID 1.
                                                                  ABK61452 standard; cDNA; 7560 BP.
Human cDNA encoding protein NOV2.
WO200216599-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DNA; 10766 BP
                                                                                                                                                                                                                                                                                                                                                                                                             WOZOCZ
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.5%;
                                                                                                                       28-FEB-2002.
(CURA-) CURAGEN CORP.
(CORT-) COR THERAPEUTICS INC.
       (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          13-MAR-2003.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                   Query Match
Best Local Similarity
RESULT 722
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 728
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Best Local Similarity
RESULT 729
                                                                                                                                                                                          Best Local Similarity RESULT 723
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RESULT 730
ID AD034038 standard;
DE Human CLA2 gene, SB
                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 724
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Best Local Similarity
RESULT 725
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Length 110000;

Length 261;

Length 371;

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Length 110000;

Length 110000,

Length 110000;

Length 110000;

Length 49561

Length 64976

Length 110000,

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Best Local Similarity 51,8%; Pred. No. 1.4e+02;
SULT 752
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                                                                                                                                                                                                                                                                                                                                                       RESULT 755
                RESULT
                                                                                                                                                                                                                                                                                                                                                                                    ACA71987 standard; DNA, 457 BP.
Human PRO polypeptide associated oligonucleotide SEQ ID NO 432.
US2002177553-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 20-MAR-2003.
(GETH ) GENENTECH INC.
10.4%; Score 28.4; DB 9; Length 457;
 Length 416;
                                                                                                                   Length 457;
                                                                                                                                                                                                                                     Length 457;
                                                                                                                                                                                                                                                                                                                                                          457;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 457;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADA24971 standard; cDNA; 457 BP.
Novel human secreted and transmembrane protein EST DNA15846.
US2003050241-A1.
13-MAR-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ACD29969 standard; cDNA; 457 BP.
Novel human secreted and transmembrane protein EST DNA15846.
US2003050240-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 457
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 457
                                                                                                                                                                                                                                                                           ACA63823 standard; cDNA; 457 BP.
Novel human secreted and transmembrane protein DNA15846.
US2002192706-A1.
                                                                                                                                                                                                                                                                                                                                                          Length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADA12632 standard; cDNA; 457 BP.
Human secreted/transmembrane polypeptide PRO218 EST #2
US2003055216-A1.
                                                                                                                                                         AAC78565 standard; cDNA; 457 BP.
Human EST DNA15846 nucleotide sequence SEQ ID NO:432.
WO200053756-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human secreted/transmembrane protein EST DNA15846.
US2003004102-A1.
                                                                                      16-SEP-1999.
(GETH) GENENTECH INC.
ery Match 10.4%; Score 28.4; DB 2;
ery Match 10.4%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                US20ve....
28-NOV-2002.
(GFH) GENENTECH INC.
(GFY) Match
10.4%; Score 28.4; DB 8;
inilarity 51.8%; Pred. No. 1.4e+02;
Score 28.4; DB 9;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                10.4%; Score 28.4; DB 3; 51.8%; Pred. No. 1.4e+02;
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(GETH ) GENENTECH INC.

(GETH ) GENENTECH INC.

10.4%; Score 28.4; DB 8;

lery Match

10.4%; Score 28.4; DB 8;

11.40+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABX92627 standard, cDNA; 457 BP.
Human PRO218 EST polynucleotide sequence #2.
US200162284-A1.
US-NOV-2002.
(GRTH) GENENTECH INC.
ery Match
still Local Similarity 51.8%; Pred. No. 1.40+
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              L. Careted/transmembrant.

2.03004102-A1.

02-JAN-2003.

PA (GETH) GENENTECH INC.
QUERY MAtch
Best Local Similarity 51.8%; Pre
RESULT 749
ID ADA24971 standard; CDNA
DE NOVOEl human Secret
PN US200305024'
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACA66368 standard; cDNA; 457 BP
                                           DNA; 457 BP.
10.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.4%;
                                                                                                                                                                                                                     (GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 745
                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
RESULT 746
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 747
       Best Local Similarity RESULT 743
                                      AAZ34255 standard;
Human EST DNA15846.
WO9946281-A2.
                                                                                                                           Best Local Similarity
RESULT 744
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RESULT 748
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                                                                                    16-SEP-199
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Query Match
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20-MAR-2003.
(GETH) GENENTECH INC.
ery Match
ery Match 10.4%; Score 28.4; DB 10; Length 457;
                                                                                                                                                                                                                                              Score 28.4; DB 10; Length 457; Pred. No. 1.4e+02;
Novel human secreted and transmembrane polypeptide cDNA #108. US2003049633-A1.
                                                                                                                                                                                                                                                                                                                                                                                              Length 457;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 457;
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                                                                                                       Length 457;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADC63804 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2003054405-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADC69028 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human EST from secreted/transmembrane protein, PRO218 US2003049684-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  PR0218
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Human EST from secreted/transmembrane protein, PRO218
US2003068648-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PR0218
                                                                                                                                                                                                                                                                                                                                                                                          Score 28.4; DB 10;
Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
                                                                                                   10.4%; Score 28.4; DB 9; 51.8%; Pred. No. 1.4e+02;
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The Addedyed standard; CDNA; 457 BP.
DE Human EST from secreted/transmembrane protein, PN US2003060406-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                         ADC44080 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein,
US2003054986-Al.
                                                                                                                                                                                                                                                                                                ADB76654 standard; cDNA, 457 BP.

Human PRO polynucleotide sequence #108.

10.01-MAY-2003.

(GETH) GENENTECH INC.

10.4%; Score 28.4

St. Local Similarity 51.8%; Pred. No.
                                                                                                                                                                      Human PRO polynucleotide sequence #108 US2003045462-A1.
                                                                                                                                                       CDNA; 457 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADC61840 standard; cDNA; 457 BP
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20-MAR-2003.
(GETH ) GENENTECH INC.
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06-MRA-2003.
(GETH) GENENTECH INC.
"Match 17:rity 51.8%;
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51.8%;
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(GETH ) GENENTECH INC.
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03-APR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 761
                                                                                                              Best Local Similarity RESULT 753
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Best Local Similarity
RESULT 754
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Best Local Similarity
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RESULT 756
                                                                                                                                                       ADB73938 standard;
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Length 457;

Length 457,

Query Match

Query Match

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U3-MAR-2003.
(GETH) GENENTECH INC.
ery Match 10.4%; Score 28.4; DB 10; Length 457;
ery Match 10.4%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ACD42788 standard; cDNA; 457 BP. Novel human secreted and transmembrane protein EST DNA15846. US2003050239-A1.
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Human EST from secreted/transmembrane protein, PRO218 #2.
US2003130181-A1.
Human EST from secreted/transmembrane protein, PRO218 #2
US2003194781-A1.
                                                                                                                                                                                                                                     Aur47/272 standard; CDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218
US2003195333-A1.
                                                                                                                                                                                                                                                                                                                                                                                         ADG53029 standard; cDNA; 457 BP.

Human EST from secreted/transmembrane protein, PR0218
10/52003216511-Al.
20-NOV-2003.
(GETH) GENENTECH INC.
10.4%; Score 28.4; DB 10; Leng
stry Match
51.8%; Pred. No. 1.4e+02;
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Human EST from secreted/transmembrane protein, PRO218
                                                                                                                   ADE17258 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218
US2003203433-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PR0218
                                                                                                                                                                                                    Score 28.4; DB 10;
Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
                                                                  Score 28.4; DB 10;
Pred. No. 1.4e+02;
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Human EST from secreted/transmembrane protein,
US2003206915-Al.
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16-OCT-2003.
(GETH ) GENENTECH INC.
10.4%;
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06-NOV2003.
(GETH ) GENENTECH INC.
10.4%;
                 US20031>-.
16-007-2003.
(GETH ) GENENTECH INC.
10.4%;
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                                                                                                                                                                    30-OCT-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             GODDARD A.
GODOWSKI P J.
GIRMALDI J C.
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FERRARA N.
FILVAROFF E.
FONG S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 776
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                                                                             Best_Local Similarity
RESULT 771
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
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Best Local Similarity
RESULT 775
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(GODO/)
(GIRM/)
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                                                                                     Length 457;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 457;
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Human EST from secreted/transmembrane protein, PRO218 #2.
US2003436-A1.
30-OCT-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADE35520 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PR0218 #2.
US2003203434-A1.
30-OCT-2003.
(GETH ) GENENTECH INC.
     ADC68153 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2
US2003069178-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                     Human EST from secreted/transmembrane protein, PR0218 #2
US2003073624-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADC42097 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2003104998-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADE49466 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2
US2003096744-A1.
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US2003203435-A1.
                                                                                                                                     ADCA1473 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2
US2003072745-A1.
                                                                                                                                                                                                                                                                       ADC67528 standard; cDNA; 457 BP.

Human EST from secreted/transmembrane protein, PR0218
US200307311-A1.

17-ARR-2003.

(GETH ) GENENTECH INC.
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(GETH) GENENTECH INC.
(GETH) MATCh 10.4%; Score 28.4; DB 10;
GETY MATCh 11arity 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                           10.4%; Score 28.4; DB 10; 51.8%; Pred. No. 1.4e+02;
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(GETH) GENENTECH INC.
ery Match imilarity 51.8%; Score 28.4; DB 10;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
                                                                                     10.4%; Score 28.4; DB 10; 51.8%; Pred. No. 1.4e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADE16634 standard; cDNA; 457 BP.
                                                                                                                                                                                                                                                                                                                                                                                                              ADC62464 standard; cDNA; 457 BP
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30-OCT-2003.
(GETH ) GENENTECH INC.
10.4%;
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17-PR-2003.
(GETH ) GENENTECH INC.
10.4%;
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 768
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Best Local Similarity
RESULT 766
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Best Local Similarity
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SULT 763
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Best Local Similarity
RESULT 765
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                                                                                       Query Match
Best Local Similarity
RESULT 762
                                                                                                                                                                                          17-APR-2003
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Query Match

Length 457;

Length 457

Length 457

Length 457

Query Match

RESULT 769 BB

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Length 457;
                                                                         Length 457;
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        Human EST from secreted/transmembrane protein, PRO218 #2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADF25492 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2003211092-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADF25593 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2003199674-A1.
                                                                                                                                                                                                                                                     ADF41447 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2003199435-A1.
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Human EST from secreted/transmembrane protein, PRO218 #2
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Human EST from secreted/transmembrane protein, PRO218 #2.
US2003195344-Al.
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Human EST from secreted/transmembrane protein, PRO218 #2.
                                                                                                                                           PR0218
                                                                                                                                                                                                                                                                                                                                                                                  ADF33126 standard; cDNA; 457 BP.

Human EST from secreted/transmembrane protein, PR0218
US2003211091-A1.
13-NOV-2003.
(GETH ) GENENTECH INC.
10.4%; Score 28.4; DB 12; Leng
st Local Similarity 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.4%; Score 28.4; DB 12; 51.8%; Pred. No. 1.4e+02;
                                                                        Score 28.4; DB 12;
Pred. No. 1.4e+02;
                                                                                                                                                                                                        Score 28.4; DB 12;
Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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Pred. No. 1.4e+02;
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                                                                                                                                       Human EST from secreted/transmembrane protein, US2003199437-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADGSO605 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein,
US2003207803-A1.
Human EST 12.
US2003199436-A1.
23-OCT-2003.
(GETH ) GENENTECH INC.
"arch 10.4%; SC
                                                                                                                         ADF27853 standard; cDNA; 457 BP.
                                                                                                                                                   US20031>>...
23-OCT-2003.
(GETH ) GENENTECH INC.
10.4%; SC
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23-OCT-2003.
(GETH ) GENENTECH INC.
10.4%; SC
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13-NOV-2003.
(GETH ) GENENTECH INC.
10.4%; SC
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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                                                                                     Best Local Similarity
RESULT 786
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RESULT 788
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Best Local Similarity
RESULT 789
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                                                                                                                                                                                                                                                                                                                                                     Score 28.4; DB 12; Length 457; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.4%; Score 28.4; DB 12; Length 457; 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                      Length 457;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 457;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 457;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 28.4; DB 12; Length 457
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                ADF40199 standard; cDNA; 457 BP.

Human EST from secreted/transmembrane protein, PR0218 #2.

US200319994-A1.

23-OCT-2003.

(GETH) GENENTECH INC.

ery Match

total Similarity 51.8%; Pred. No. 1.48+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADF40823 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2003199021-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADF45995 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2
US2003195148-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADF23767 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human EST from secreted/transmembrane protein, PRO218 #2
US2003204055-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human EST from secreted/transmembrane protein, PRO218 #2
US2003194780-A1.
                                                                                                                                                                                                                                                                                    Human EST from secreted/transmembrane protein, PRO218 US2003195345-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 28.4; DB 12;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                        10.4%; Score 28.4; DB 12;
51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.4%; Score 28.4; DB 12; 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ID ADF27217 standard; cDNA; 457 BP.
                                                                                                                                                                                                                                                                    ADF61507 standard; cDNA; 457 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADF24391 standard; cDNA; 457 BP.
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                                                                                                                                                                                                                                                                                                US20031500-
16-OCT-2003.
(GETH ) GENENTECH INC.
Watch 10.4%; Sc
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16-OCT-2003.
(GETH ) GENENTECH INC.
****Ch 10.4%; SC
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51.8%;
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51.8%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Lery Match
Best Local Similarity PRESULT 781
ID ADF24391 standaring DE Human EST f-PN US20032.
PD 30-C
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                           2003.
2003.
21H ) GENENTECH 1.
2ry Match
Best Local Similarity 2.
RESULT 780
ID ADF4595 stand?
DE Human EST f
PN US20031
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (GETH ) GENENTECH INC.
                                                                                  PAN J.
PAONI N F.
ROY M A.
SHELTON D L.
STEWART T A.
                                                                                                                                                                    (TUMA/) TUMAS D.
(WILL/) WILLIAMS P M.
     GURNEY A L.
HILLAN K J.
KLJAVIN I J.
                                                                                                                                                                                                                                  Best Local Similarity RESULT 778
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 785
                                                     KUO S S.
NAPIER M A.
                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 779
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 783
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Best Local Similarity
RESULT 784
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Best Local Similarity
RESULT 782
                                                                                                                                                                                                   (WOOD/) WOOD W I.
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                                                                                                                                                                                                                        Query Match
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                      (HILL/) | (KLJA/) | (KLJA/) | (KUOS/) | (NAPI/) | (PANJ/) | (PAON/) | (ROYM/) | (SHEL/) | (STEW/) | (STEW/) | (
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        PA
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Query Match

Query Match

Query Match

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10.4%; Score 28.4; DB 10; Length 771; 56.4%; Pred. No. 1.7e+02;
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                                                  Length 457;
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                                                                                                                                                                                                                       Length 651;
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Human immune system associated gene SEQ ID NO: 2306
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human breast cancer expressed polynucleotide 16645.
WO200151628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         RESULT 809
ID ADK62643 standard; DNA; 771 BP.
DE Disease traating protein complex-derived gene #447.
PN BP1338608-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                       AALIS342 standard; cDNA; 732 BP.
Human breast cancer expressed polynucleotide 7799, w0200151528-A2.
19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                  DB 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.4%; Score 28.4; DB 10; 56.4%; Pred. No. 1.7e+02;
                                                                                                                                                                                                           Luciy Match 10.4%; Score 28.4; DB 6; Best Local Similarity 51.6%; Pred. No. 1.6e+02; RESULT 805
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.4%; Score 28.4; DB 4;
real Similarity 52.5%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 28.4; DB 6;
Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                 10.4%; Score 28.4; DB 6; 51.6%; Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.4%; Score 28.4; DB 4; 52.5%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          28.4; DB 3;
No. 1.9e+02;
01-APR-2004.
(GETH ) GENENTECH INC.
ery Match 10.4%; Score 28.4; DB 12;
ery Match 10.4%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAX20003 standard; DNA; 1069 BP.
Enterococcus faecalis EF002 gene fragment.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human secreted protein cDNA sequence WO200055199-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.4%; Score 50.0%; Pred.
                                                                                                    ABQ69334 standard; DNA; 651 BP.
Listeria innocua DNA sequence #773.
W0200228991-A2.
11-APR-2002.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                                                                                                ABO67667 standard; DNA; 705 BP.
Listeria innocua DNA sequence #469.
WO200228891-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  CDNA; 746 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ACC61006 standard; DNA; 771 BP.
Gene sequence #SEQ ID 794.
EP1258494-A1.
                                                                                                                                                                                                                                                                                                                                                          (INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    MCACAL 2002.
(BDIG-) EPIGENOMICS AG.
erv Match 10.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    21-SEP-2000.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                20-NOV-2002.
(CELL-) CELLZOME AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (CELL-) CELLZOME AG.
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Best Local Similarity
RESULT 808
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
RESULT 806
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
RESULT 807
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAL24188 standard;
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                                                                                                                                                                                                                                                                                                                                                                                                     Ouery Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                RESULT 812
                                                                                                                             ADH25654 standard; cDNA; 457 BP.
Human neurotrimin homologue related nucleotide sequence SEQ ID NO:432.
EP1386931-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 457;
                                                                      Length 457;
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Best Local Similarity 51.8%; Pred. No. 1.4e+02;
RESULT 802
DE ADM17431 standard; CDNA; 457 BP.
DE Human BST from secreted/transmembrane protein, PRO218 #2.
PN US2004048332-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADL07265 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2.
US2004063921-A1.
                                                                                                                          ACK51853 standard; cDNA; 457 BP.

Human EST from secreted/transmembrane protein, PRO218 #2.

US2003215908-A1.

20-NOV-2003.

(GETH ) GENENTECH INC.

10.4%; Score 28.4; DB 12; Length

st Local Similarity 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADG59173 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218 #2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human BST from secreted/transmembrane protein, PRO218 #2 US2004006219-A1.
                                                                                                                                                                                                                                                                                                    Human BST from secreted/transmembrane protein, PRO218 US2003216305-A1. 20-NOV-2003. (GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                       ADG48733 standard; cDNA; 457 BP.
Human EST from secreted/transmembrane protein, PRO218
US2003216560-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADG51229 standard; cDNA; 457 BP.

Human EST from secreted/transmembrane protein, PRO218
US2004005312-A1.

(GB-JAN-2004.

(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
ery Match 10.4%; Score 28.4; DB 12;
ery Match 51.8%; Pred. No. 1.4e+02;
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08-JAN-2004.

(GETH ) GENENTECH INC.

10.4%; Score 28.4; DB 12;

10.4%; Score 28.4; DB 12;

10.4%; Pred. No. 1.4e+02;
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(GETH ) GENENTECH INC.

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10.4%; Score 28.4; DB 12;

(ery Match

1 cimilarity 51.8%; Pred. No. 1.4e+02;
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20-NOV-2003.
(GETH) GENENTECH INC.
ery Match
10.4%; Score 28.4; DB 12;
ery Match
51.8%; Pred. No. 1.4e+02;
                                                                      10.4%; Score 28.4; DB 12; 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                   10.4%; Score 28.4; DB 12; 51.8%; Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                      ADG49357 standard; cDNA; 457 BP
                                  20-NOV-2003.
(GETH ) GENENTECH INC.
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RESULT 796
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Best Local Similarity
RESULT 798
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Best Local Similarity
SULT 799
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                                                                                      Best Local Similarity RESULT 795
                                                                                                                                                                                                                                                                                                                                                                                                 Best_Local Similarity RESULT 797
                US2003215905-A1.
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08-APR-2004
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAX20002 standard; DNA; 1161 BP.
Enterococcus faecalis gene BF002.
WOSB5554-A2.
12-NO-1998.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
st Local Similarity 62.9%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.4%; Score 28.4; DB 10; Length 1296; 62.9%; Pred. No. 2.1e+02;
                                                            Length 1069;
                                                                                                                                                                                                          Query Match 10.4%; Score 28.4; DB 6; Length 1069; Best Local Similarity 62.9%; Pred. No. 1.9e+02; RESULT 814
                                                                                                                                                                                                                                                                                                                                                               Query Match 10.4%; Score 28.4; DB 8; Length 1069; Best Local Similarity 62.9%; Pred. No. 1.9e+02; RESULT 815
                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.5EP-2002.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match
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10.4%; Score 28.4; DB 8; Length 1069;
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ry Match
10.4%; Score 28.4; DB 8; Length 1161;
t Local Similarity 62.9%; Pred. No. 2e+02;
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        T2-NOV-1998.
(HUMA-) HUMAN GENOME SCI INC.
10.4%; Score 28.4; DB 2;
lery Match
10.4%; Pred. No. 1.9e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                       ABX61558 standard; DNA; 1069 BP.
Enterococcus faecalis EF040 polynucleotide #4.
US6448043-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABX61557 standard; DNA; 1161 BP.
Enterococcus faecalis EF040 polynucleotide #3
US6448043-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADH85017 standard; DNA; 1296 BP.
Enterococcus faecalis polynucleotide #2902.
US6617156-B1.
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Human GPCR gene SEQ ID NO:21.
                                                                                                                                                                                                                                                               ACA87952 standard, DNA; 1069 BP. E. faecalis novel gene #4. 025003017495-A1. 23-JAN-2003. (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AEN97987 standard, DNA; 1161 BP. E faecalis EF002 gene. US2002045737-A1. 18-APR.2002. (HUMA-) HUMAN GENOME SCI INC.
                                                                                                        ABN97988 standard, DNA; 1069 BP.
E faccalis EF002 gene fragment.
18-APR-2002.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ACA87951 standard; DNA; 1161 BP.
E. faecalis novel gene #3.
US2003017495-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10-SEP-2002.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (DOUC/) DOUCETTE-STAMM L A. (BUSH/) BUSH D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity 6
RESULT 816
ID AAX20002 standard; DNA
DE Enterococcus faecalis
PN W99850554-A2.
PD 12-NOV-1998
PA (HUMA-) HUMAN GENOME S
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity 6
RESULT 819
ID ARX61557 standard; DNA
DE Enterococcus faecalis
PD 10-SEP-2002.
PA (HUMA-) HUMAN GENOME S
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Lery Match
Best Local Similarity to RESULT 821
ID ADC85568
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity RESULT 817
                                                      Query Match
Best Local Similarity
RESULT 813
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 820
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WO9850554-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            09-SEP-2003
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PD 02-JAN-2003.

PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.

PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.

Query Match

Best Local Similarity 51.6%; Pred. No. 2.1e+02;

RESULT 822
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.4%; Score 28.4; DB 12; Length 2074; 60.3%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ID ABX34524 standard; cDNA; 2311 BP.

DE Human mddt cDNA SEQ ID 85.

PN WO200279449-A2.

PD 10-OCT-2002.

PA (INCY-) INCYTE GENOMICS INC.

Query Match 10.4%; Score 28.4; DB 8; Length 2311;

Best Local Similarity 46.1%; Pred. No. 2.56+02;
                                                                                                                                                                                                                                                             Length 1451;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 2280;
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(INCY-) INCYPE GENOMICS INC.
(ery Match
(ery Match 10.4%; Score 28.4; DB 6; Length 2281;
(ery Match 16.1%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.4%; Score 28.4; DB 4; Length 2750; 60.3%; Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAS31144 standard; cDNA; 2281 BP.
Human diagnostic and therapeutic polynucleotide (DITHP) #159
WO200162927-A2.
                                                                                                                                                                                                                                                                                                        ADVS7300 standard; cDNA; 2074 BP.
DNA encoding Kidney development associated protein segid 67.
US2044068763-A1.
                                                                                                                                              AAF88594 standard; cDNA, 1451 BP.

Human GCREC-15 cDNA INCYTE ID 7475271CD1 SEQ ID 6
W02002633004-A2.
W1047-002.
(INCY-) INCYTE GENOMICS INC.
ery Match

Local Similarity 51.6%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.4%; Score 28.4; DB 2; 46.1%; Pred. No. 2.5e+02;
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Query Match
10.4%; Score 28.4; DB 6;
Best Local Similarity 46.1%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL99755 standard; cDNA; 2280 BP.
Human secretory polynucleotide (sptm) 10.
WO200220756-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DE Human cDNA sequence SEQ ID NO:18992.
PN EP1074617-A2.
D 07-FEB-2001.
A (HELI-) HELIX RES INST.
Best 7.-.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAV74265 standard; cDNA; 2268 BP.
Human heart muscle specific cDNA #3.
WO9856907-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABS70446 standard; cDNA; 2281 BP. Human bone remodelling gene #103. US6426186-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              30-AUG-2001.
(INCY-) INCYTE GENOMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                            (HOPK/) HOPKINS N.
(GOLL/) GOLLING G.
(AMST/) AMSTERDAM A.
(SUNZ/) SUN Z.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               17-DEC-1998.
(MEDI-) MEDIGENE AG.
                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 823
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 824
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity
RESULT 825
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Length 7393;

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Score 28.4; DB 9;
Pred. No. 3.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                    WO20027,2...
03-OCT-2002.
(EPIG-) EPIGENOMICS AG.
10.4%; Sc
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         WOZUGECO
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Wolvot-2
01-NOV-2001.
(EPIG-) EPIGENOMICS AG.
10.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             09-AUG-2001. (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.4%;
               10.4%;
                                                                                                                                                                                                                                                                                                           10.4%;
(DRMA/) DRMANAC R T.
                                                                                                                                                                                                                                                         (KUNS/) KUNSCH C A.
(DILL/) DILLON P J.
(BARA/) BARASH S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 844
               Query Match
Best Local Similarity
RESULT 839
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ID ABL92304 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAK83881 standard;
                                                                                                                                                                                                                                                                                                                           Local Similarity
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                                                                                                                                                                                                                                                                                                             Query Match
                  ADG3304B standard; DNA; 2750 BP.
Whaman DNA differentially expressed in patients with SLE SeqID372.
WC2003090694-A2.
06-NOV-2003.
                                                                                                                                        ABL25494 standard; DNA; 3371 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 27955.
WO200171042-A2.
                                                                                                Length 2750
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AG.
10.4%; Score 28.4; DB 6; Length 6640;
49.3%; Pred. No. 3.7e+02;
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                                                                                                                                                                                                                                Length 3371;
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Drosophila melanogaster genomic polynucleotide SEQ ID
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABL33657 standard; DNA; 6640 BP.
Human immune system associated gene SEQ ID NO: 1630.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DNA; 5962 BP.
n associated gene SEQ ID NO: 1259
                                                                                (EXPR-) EXPRESSION DIAGNOSTICS INC.
ry Match 10.4%; Score 28.4; DB 10;
t Local Similarity 60.3%; Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                      09-MRR-1994.

(FARH ) HOECHST JAPAN LTD.

(FARH ) HOECHST JAPAN LTD.

ery Match

o'milarity 54.9%; Pred. No. 3e+02;
                                                                                                                                                                                                                                10.4%; Score 28.4; DB 4;
45.0%; Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.4; DB 6;
Pred. No. 3.6e+02;
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Pred. No. 3.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADQ98702 standard; cDNA; 7393 BP.
DNA encoding human GPCR-like protein segid 372
US6569662-B1.
                                                                                                                                                                                                                                                            Sequence of human OSF-4-1 cDNA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAIS8487 standard; cDNA; 7393 BP.
Human polynuclectide SEQ ID NO 690
WO200153312-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ____sv_ standard; cDNA; 7393 BP.
Novel human cDNA SEQ ID NO 372.
US2003104529-A1.
05-TIM...o.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AG.
10.4%;
47.7%;
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52.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABL33286 standard; DN
Human immune system as
WOZ00209298-A2.
03-JAN-2002.
(EPIG-) EPIGENOMICS A
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(EPIG-) EPIGENOMICS
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Best Local Similarity
RESULT 838
                                                                                                                                                                                           27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                         27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity RESULT 835
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RESULT 836
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RESULT 832
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Best Local Similarity
RESULT 834
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Best Local Similarity
RESULT 833
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Best Local Similarity
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(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ZHOU P.
TANG Y T.
LIU C.
ASUNDI V.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    26-JUL-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               05-JUN-2003
(ZHOU/) ZHO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (TANG/)
(LIUC/)
(ASUN/)
                                                                                                   Query Match
                                                                                                                                                                                                                                 Query Match
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RESULT 837
     RESULT 830
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ABZ09978 standard, DNA, 7809 BP.
Haematopoietic cell proliferation disorder related DNA sequence #118.
W0200277272-A2.
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WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        14396
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 10318;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 10696;
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                                                                                                              Length 7574;
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27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABK39952 standard; DNA; 17703 BP.
Human chemically pretreated gene sequence #17 strand 1.
WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL54340 standard; DNA; 10696 BP. Chemically treated apoptosis gene complementary to WO200177164-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human immune system associated gene SEQ ID NO: 292 WO200200928-A2.
                           contig SEQ ID NO:152
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 18-007-2001.

18-007-2001.

(EPIG-) EPIGENOMICS AG.

(EPIG-) BPIGENOMICS AG.

10.4%; Score 28.4; DB 6;

(ery Match 1arity 48.7%; Pred. No. 4.4e+02;
AAX13089 standard; DNA; 7574 BP.
Enterococcus faecalis genome contig SEQ ID NO:15
W09850555-A2.
12-NOV-1998.
(HUMA) HUMAN GENOME SCI INC.
ery Match
st Local Similarity 62.9%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 28.4; DB 4;
Pred. No. 4.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 28.4; DB 6;
Pred. No. 4.8e+02;
                                                                                                                                                                                                                                                                                                                                      Score 28.4; DB 6;
Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.4%; Score 28.4; DB 4; 47.7%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 28.4; DB 8;
Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 28.4; DB 6;
Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADDY2004 Standard; DNA; 13919 BP. Chemically treated DNA repair gene fragment#57. WO200181622-A2.
                                                                                                                                                                              ABS98884 standard; DNA; 7574 BP.
Enterococcus faecalis contig sequence #152.
US2002120116-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABL06638 standard; cDNA; 10318 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DNA; 10696 BP
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Score 28.4; DB Pred. No. 1e+03;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.4%; Score 28.4; DB 51.6%; Pred. No. 1e+03;
                                                                                   Human MAP kinase MAPKI0 (JNK3) gDNA. W02004047623-A2. I0-U0V-2004.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Listeria innocua DNA sequence #684
WO200228891-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADC85227 standard; DNA; 96599 BP. Mouse Irf2 genomic sequence. W02003045230-AZ. 05-JUN-2003.
                                                                                                                                                                                                                                                                                                                                                                                                   ADB72485 standard; DNA; 96599 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DNA; 3011208
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI
                                                                                                                                                                                                                                                                                                        N-2-0-2003.
(SAGR-) SAGRES DISCOVERY.
Lery Match 10.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.4%;
                                                                                                                                                                                                                  10.4%;
50.7%;
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                                                                                                                                                                 10.4%;
52.5%;
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76.1%;
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51.6%;
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51.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.4%;
51.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                  Mouse Irf2 gene.
WO2003008583-A2.
30-JAN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   15-APR-2004.
(MORR/) MORRIS D W.
(ENGE/) ENGELHARD E K.
               27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                Best Local Similarity RESULT 857
                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 860
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 862
                                                                                                                                                               Query Match
Best Local Similarity
RESULT 858
                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 859
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 861
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 863
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 866
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 869
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
RESULT 868
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABQ69245 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
 WO200171042-A2,
                                                    Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 864
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human membrane spanning 4-domain family, subfamily A genomic sequence. WO200262946-A2.
                                                                            AAL37510 standard; DNA; 31314 BP.
Human musculoskeletal system related polynucleotide SEQ ID NO 3875.
WO200155367-A1.
                                                                                                                                                                                                                                                                                                                                                                            AD031248 standard; DNA; 31314 BP.

Human musculoskeletal system-associated genomic DNA - SEQ ID 3875.
US2004009488-A1.
15-JAN-2004.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
st Local Similarity 53.6%; Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL11838 standard; cDNA; 49380 BP. Drosophila melanogaster expressed polynucleotide SEQ ID NO 29996.
                                                                                                                                                                                                            ABX60498 standard; cDNA; 31314 BP.
cDNA encoding novel human musculoskeletal system antigen #2842.
US2002147140-A1.
                                10.4%; Score 28.4; DB 6; Length 17703; 47.3%; Pred. No. 5.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL98269 standard; DNA; 31949 BP.
Human testicular antigen encoding DNA fragment SEQ ID NO: 2921
WO200155317-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL98270 standard; DNA; 32186 BP.
Human testicular antigen encoding DNA fragment SEQ ID NO: 2922
WO200155317-A2.
                                                                                                                                                               Length 31314;
                                                                                                                                                                                                                                                                                                                                 Length 31314;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               PD 02-AUG-2001.

Query Match Thuman GENOME SCI INC.

Query Match Thuman GENOME SCI INC.

Query Match Thuman GENOME SCI INC.

10.4%; Score 28.4; DB 4; Length 31949; Best Local Similarity 52.5%; Pred. No. 6.5e+02; RESULT 853

ID AlloS411 standard; DNA; 32186 BP.

DE Human reproductive system related antigen DNA SEQ ID NO: 8099.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (HUMA-) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 52.5%; Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
Ery Match
ery Match
From Score 28.4; DB 4; Length 32186;
From Score 28.4; DB 4; Length 32186;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AALOS410 standard; DNA; 31949 BP.
Human reproductive system related antigen DNA SEQ ID NO: 8098
WO200155320-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 32186;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 38736;
                                                                                                                      Score 28.4; DB 8;
Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.4%; Score 28.4; DB 4; 52.5%; Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.4%; Score 28.4; DB 6; 49.3%; Pred. No. 7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABQ99652 standard; DNA; 38736 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                 10.4%;
            (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                        JAN.

JON4.

JON4.

JON4.

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BEST LOCAL SIMILARITY

RESULT 851

ID AALOS410 stand?

DE Human repro

PN WO2001.

PD 02.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           J-A2.

2001.

207.

-2ry Match

Best Local Similarity to RESULT 854

ID ABL98270 stand?

D ABL98270 stand?

PN WC2001.

PN WC2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        JU-A2.

JU-A2.

JU-A2.

JUA-) HUMAN GENON.

JUA-) HUMAN GENON.

Best Local Similarity RESULT 852

ID ABL98269 stand?

DE Human test?

PN W02001=7

PD 02-7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
RESULT 855
ID A8699652 standard; DN
DE Human membrane spanni
PN W0200262946-A2.
PD 15-AUG-2002.
PA (UYDU-) UNIV DUKE.
                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
REGULT 850
ID AbJ31248 standard; DP
DE Human musculoskeletal
PN US2004009488-A1.
PD I5-JAN CBNOME
PA (HUMA-) HUMAN GENOME
                                                                                                                                                                                                                                                             10-OCT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                         Best Local Similarity RESULT 848
                                                                                                                                                              Query Match
Best Local Similarity
RESULT 849
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 856
10-JAN-2002
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Score 28.4; DB 12; Length 76500; Pred. No. 8.9e+02;
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                                                                                                                                                                                                                                                                                                                     Length 96599;
Length 49380;
                                                                                                                                                                             Length 95269;
                                                                                                                                                                                                                            ADA02747 standard; DNA; 96599 BP.
Mouse Irf2 carcinoma associated gene, SEQ ID NO:1265.
WO2003057146-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 28.4; DB 10;
Pred. No. 9.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.4%; Score 28.4; DB 10; 60.3%; Pred. No. 9.7e+02;
10.4%; Score 28.4; DB 4; 50.0%; Pred. No. 7.6e+02;
                                                                                                                                                                                                                                                                                                                     Score 28.4; DB 9;
Pred. No. 9.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADM74342 standard, DNA; 96599 BP.
Murine carcinoma associated (CA) nucleic acid #7
US2004072154-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28.4; DB 12;
Pred. No. 9.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 28.4; DB 6;
Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.4; DB 6;
Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 6;
                                                                                                                                                                           Score 28.4; DB 6;
Pred. No. 9.6e+02;
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Pred. No. 1e+03;
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ADB92114 standard; DNA; 183610 BP.

Human MDR1 related DNA sequence SEQ ID NO:664.

MOX003013535-A2.

20-FEB-2003.

(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.

ET Match

10.4%; Score 28.4; DB 10; Length 183610; st Local Similarity 50.7%; Pred. No. 1.2e+03;
   Score 28.4; DB 10; Length 183610;
Pred. No. 1.2e+03;
                                                                                                                                                  Length 183610;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.4; DB 12; Length 326014; Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 326014;
                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT BESONT BESONT BESONT BE.

ID ABK892956 standard; DNA; 326014 BE.

DE Human gene for novel serine/threonine serine kinase.

Query Match

10.4%; Score 28.4; DB 6; Length 326014;

Part Loral Similarity 51.6%; Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                               ID ABK89296 standard; DNA; 326014 BP.

DB Human gene for novel serine/threonine serine kinase.

Query Match 10.4%; Score 28.4; DB 6; Length 326014;

Best Local Similarity 53.6%; Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 304;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 233;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          9
                                                                                                       Score 28.4; DB 12;
Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28.2; DB 10;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human secreted protein 5' EST, SEQ ID NO: 20662.
EP1033401-A2.
06-SEP-2000.
(GEST ) GENSET.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AATT5253 standard; DNA; 233 BP.
Nucleotide sequence encoding human RAD50 intron
W0972784-A2.
31-JUL-1997.
(GENELABS TECHNOLOGIES INC.
ery Match
st Local Similarity 53.1%; Pred. No. 1.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 28.2; DB 6;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.2; DB 3;
Pred. No. 1.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAC09513 standard; cDNA; 380 BP.
Human secreted protein 5' EST, SEQ ID NO: 13588
                                                                          DNA sequence SEQ ID NO:664
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABX88534 standard; cDNA; 304 BP.
Corn ear-derived polynucleotide (cpd) #6994.
US6476212-B1.
                                                          DNA; 183610 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADQ94991 standard; DNA; 326014 BP.
Filman Kinase genomic DNA.
Ouery March
Best Local Similarity 53.6%; Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DNA; 326014 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (GENO-) GENOME THERAPEUTICS CORP
   10.4%;
50.7%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.3%;
52.1%;
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rry Match 10.3%;

it Local Similarity 55.7%;
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51.6%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ID ADQ94981 standard; DNA; 3:
DE Human Kinase genomic DNA.
Query Match
Best Local Similarity 51.67
RESULT 888
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
RESULT 889
   Query Match
Best Local Similarity
RESULT 882
                                                                          Human MDR1 related WO2003013537-A2.
                                                                                                                                                                     Best Local Similarity
                                                                                                                                                                                                                                                                                                                     Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
RESULT 891
                                                        ADB96923 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    05-NOV-200
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                                                                                                                                                                                                                                                                             AAH44801 standard; DNA; 112190 BP.

Human GPCR protein KAT06734L DNA containing exons 3 to 9 SEQ ID NO:16.
JP2001245666-A.

11.SEP-2001.

(KYOW ) KYOWA HAKKO KOGYO KK.

10.4%; Score 28.4; DB 4; Length 112190;

SEL Local Similarity 52.5%; Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                        SeqID 1.
129017;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADP65634 standard; DNA; 137908 BP.
Human sequence from clone 914P14 on chromosome Xq23 Contains DNA
WO2003072827-Al.
                                                                                                                 Length 110000
                                                                                                                                                                       Length 110000;
                                                                                                                                                                                                                            Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 137908;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.4%; Score 28.4; DB 8; Length 183610; 50.7%; Pred. No. 1.2e+03;
   Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 183610;
                                                      Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 168174;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 168273;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    63
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ACF62736 standard; DNA; 183610 BP.
Cancer based on CYP3A5 related polynucleotide SEQ ID NO:664
WO2003013534-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                 ADP84158 standard, DNA; 129017 BP. Human AST-1 locus DNA representing part of the GPRA generry Match 50.00 10.44; Score 28.4; DB 12; Length st Local Similarity 52.54; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABT11173 standard; DNA; 168174 BP.
Human 5-lipoxygenase gene related DNA sequence SEQ ID No
WO200262825-A2.
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Human 5-lipoxygenase gene related DNA sequence SEQ ID
WO200262825-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADB20851 standard; DNA; 183610 BP.
MRP1 based cancer related nucleic acid SEQ ID NO:664.
WO2003013533-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             O4-SEP-2003.
(CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT.
ery Match
10.4%; Score 28.4; DB 11;
                                                                                                                                                                                                                        Score 28.4; DB 10;
Pred. No. 1e+03;
Score 28.4; DB 8;
Pred. No. 1e+03;
                                                                                                             Score 28.4; DB 10;
Pred. No. 1e+03;
                                                                                                                                                                     Score 28.4; DB 10;
Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         WOLUDGOOD.

15-AUG-2002.
(MILL-) MILLENNIUM PHARM INC.
(MILL-) MILLENNIUM PARM INC.
ery Match
10.4%; Score 28.4; DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  20-EBB-2003.
(BPID-) EPIDAUROS BIOTECHNOLOGIE AG.
ery Match
10.4%; Score 28.4; DB 8;
                                                          DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.4%; Score 28.4; DB 6; 56.4%; Pred. No. 1.2e+03;
                                                        Score 28.4; DB Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADB87940 standard; DNA; 183610 BP.
Human UGT1Al gene sequence SEQ ID NO:664
WO2003013536-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (BPID-) EPIDAUROS BIOTECHNOLOGIE AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            20-FEB-2003. (EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     15-AUG-2002.
(MILL-) MILLENNIUM PHARM INC.
                                                                                                               10.4%;
51.1%;
                                                                                                                                                                                                                          10.4%;
56.4%;
   10.4%;
51.6%;
                                                                                                                                                                     10.48;
                                                                                                                                                                                     56.48;
                                                        10.4%;
                                                                            50.08;
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Best Local Similarity
RESULT 876
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                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity
RESULT 875
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 879
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RESULT 880
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity RESULT 878
                                                                                                                                                                                                                        Query Match
Best Local Similarity
RESULT 874
 Query Match
Best Local Similarity
                                                                                                               Query Match
Best Local Similarity
RESULT 872
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                                                                          Local Similarity
                                                                                                                                                                                       Best Local Similarity
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                                                        Query Match
                                                                                                                                                                     Query Match
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                     Best Loc
RESULT 870
                                                                                               871
                                                                          Best
RESULT
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Prokaryotic essential gene #21209
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Prokaryotic essential gene #10011 WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACA28354 standard; DNA; 1191 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ACA39552 standard; DNA; 996
                                                                                                                                                                                                                                                                                                                                                                                                         PAGE A.
MATHEW A V.
LEDFORD B L.
WOESSNER J P.
HAAS W D.
GARCIA C A.
KRICKER M.
SLATER T.
DAVIS K R.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ...URB/) HOFFMAN N...URB/) HURBAN P...uery Match
Best Local Similarity
RESULT 902
ID ACA39552 stand?
DE Prokaryotic
PN W020027*
PD 03-**
              PAGE A.
MATHEW A V.
LEDFORD B L.
WOESSNER J P.
                                                                                                                                                                                                                                                                                                                                   HAMILTON C M.
                                                                          HAAS W D.
GARCIA C A.
KRICKER M.
SLATER T.
 RAMEAKA J G.
                                                                                                                                                                                                                                                                                                                                                                                              RAMEAKA J G.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 905
                                                                                                                                    DAVIS K R.
ALLEN K.
HOFFMAN N.
                                                                                                                                                                                                                                                                                                                                                                RAINES T M.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 904
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 903
                                                                                                                                    (DAVI/) DAVIS K R. (ALLE/) ALLEN K. (HOFF/) HOFFMAN N. (HURB/) HURBAN P.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                              Local Similarity
                                                                                                                                                                                                                                                                                                      GORLACH J.
                                                                                                                                                                                                                                                                                                                                                     PRICE J L.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
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                                                                                                                                                                                                   Query Match
                                                        (WOES/)
(HAAS/)
(GARC/)
(KRIC/)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (GARC/)
(KRIC/)
(SLAT/)
(DAVI/)
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(YUYY/)
 (RAME/)
                                                                                                                                                                                                                                                                                                                                    (HAMI/)
                                                                                                                                                                                                                                                                                                                      (ANYY/)
                                                                                                                                                                                                                                                                                                                                                                                                (RAME/)
                                                                                                                                                                                                                                                                                                                                                                                                           (PAGE/)
                 (PAGE/
                                                                                                                                                                                                                                                                                                                                                                                                                                                          WOES/
                                             LEDF,
                                                                                                                                                                                                               Best Loc
RESULT 901
                                                                                                                                                                                                                                                                                                                                                                                                                                           LEDF
                                                                                                                                                                                                                                               AEQ38867 standard; DNA; 679 BP.
Oligonucleotide for detecting cytosine methylation SEQ ID NO 25458.
WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AEQ38866 standard; DNA; 679 BP. Oligonucleotide for detecting cytosine methylation SEQ ID NO 25457. WC200218632-A2. 07-MAR-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAH92502 standard; DNA; 700 BP.
Human inflammatory bowel disease related gene fragment IGR1202a
WO200142511-A2.
                                                                                                                                                                                                                                                     MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.3%; Score 28.2; DB 5; Length 496;
                                                                                                                                                                  Length 392;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               MS-APR-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
10.3%; Score 28.2; DB 4; Length 631;
ery Match
11.2%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               WHED ) WHITEHEAD INST BIOMEDICAL RES.
(ELLI-) ELLIPSIS BIOTHERAPEUTICS CORP.

10.3%; Score 28.2; DB 4; Length 700; st Local Similarity 53.1%; Pred. No. 1.9e+02;
                                               Length 380;
                                                                                                                                                                                                                                                                                                                                                                                                             Length 516;
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1 (EPIG-) EPIGENOMICS AG.
Query Match
10.3%; Score 28.2; DB 6; Length 679;
neet focal Similarity 52.6%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.3%; Score 28.2; DB 6; Length 679 52.6%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAH33609 standard; cDNA; 631 BP.
Human colon cancer antigen encoding cDNA SEQ ID NO:665.
WO200122920-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Arabidopsis thaliana polynucleotide SEQ ID NO 214 US2002059663-A1.
                                            10.3%; Score 28.2; DB 3; 51.2%; Pred. No. 1.5e+02;
                                                                                                                                    PD 11-APR-2002.

PA (GENE-) GENE LOGIC INC.

QUERY MATCH

10.3%; Score 28.2; DB 6;

Best Local Similarity 53.1%; Pred. No. 1.5e+02;

RESULT 894
                                                                                                                                                                                                                                                                                                                                                                          13-00-2002.

(OXFO-) OXFORD BIOMEDICA UK LTD.

(OXFO-) OXFORD BIOMEDICA UK LTD.

10.3%; Score 28.2; DB 6;

ery Match 1.7e+02;

cry mailarity 52.1%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                   ABV77972 standard; DNA; 516 BP.
Hypoxia-repressed protein coding sequence #40.
WO200246465-A2.
                                                                                                                                                                                                           ABV46766 standard; cDNA; 496 BP.
Human prostate expression marker cDNA 46757.
WC200160860-A2.
                                                                                        ABN94868 standard; DNA; 392 BP.
Gene #1366 used to diagnose liver cancer.
WO200229103-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABQ65637 standard; DNA; 709 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         07-MAR-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                .ide
.32-A2.
.21-A2.
.zry Match
Best Local Similarity 5.
RESULT 899
ID AAH92502 standar?
DE Human inflamm
PN WC2001425*
PD 14-JUW
PA (WITT)
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2002.
And Best Local Similarity RESULT 898
ID ARQ38866 standa DE Oligonucler'
PN W020021
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Best Local Similarity
RESULT 897
ID A6038867 standard; DN
DE Oligonuclectide for d
PN WO200218632-A2.
PD 07-MAR-2002.
PA (EPIG-) EPIGENOMICS AV
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HAMILTON C M.
                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
RESULT 896
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 PRICE J L.
RAINES T M.
YU Y.
                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 895
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 900
                                                        Best Local Similarity RESULT 893
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (GORL/) GORLACH J. (ANYY/) AN Y.
EP1033401-A2.
06-SEP-2000.
(GEST ) GENSET.
                                                                                                                                                                                                                                                                                          Query Match
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                                             Query Match
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(RAIN/)
(YUYY/)
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10.3%; Score 28.2; DB 6; Length 1409; 53.1%; Pred. No. 2.4e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 1191;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      MACKUTT-2002.
(BLITT-) BLITRA PHARM INC.
(ELITT-) ELITRA PHARM INC.
ery Match
10.3%; Score 28.2; DB 8; Length 996;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.3%; Score 28.2; DB 6; Length 856; 48.0%; Pred. No. 2e+02;
                                                                       ABN98845 standard; DNA; 856 BP.
Arabidopsis thaliana expressed polynucleotide SEQ ID NO 613.
US2002023231-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABK49115 standard; cDNA; 1409 BP.
Human cDNA encoding transcription factor LCR-F19.02
WO200220583-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAH99669 standard; cDNA; 1442 BP.

Human protein encoding cDNA sequence SEQ ID NO:504, N W0200153455-A2.

26-JUL-2001.

A (HYSE-) HYSEQ INC.

Query Match.
10.3%; Score 28.2; DB 6; 53.1%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         WCACCT.

03.07.

(BLITT-) ELITRA PHARM INC.

(ELIT-) ELITRA PHARM INC.

10.3%; Score 28.2; DB 8;

(ery Match

' rimilarity 47.0%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             14-MAR-2002.
(SHAN-) SHANGHAI BIOWINDOW GENE DEV INC.
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10.3%;
53.1%;
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51.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.3%;
                                                                                                                                                                                                                                                                 (ELIT-) ELITRA PHARM INC.
ry Match
t Local Similarity 47.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAVOSSSI standard; cDNA;
Human TRAF-2 kinase gene.
WO9801541-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    15-AUG-2002.
(UYMA-) UNIV MAASTRICHT.
                                                                                                                                                                                                                                                                                                                                                       LAG-A2.

LAZ-2004.

LAZEL-) EXELIXIS IN

LOGAL Similarity ESSULT 918

ID ADN04785 stand*

DE Antipsoriat*

PN W020040*

PD 08-
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(EXEL-) EXELIXIS INC.
21-NOV-2002.
A (CORI-) CORIXA CORP.
Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                15-JAN-1998.
(TULA-) TULARIK INC.
                                                                                                                                           (CORI-) CORIXA CORP.
                                       Best_Local_Similarity
RESULT 915
                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 917
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity RESULT 919
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best_Local Similarity RESULT 920
                                                                                                                                                                     Best Local Similarity RESULT 916
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Local Similarity
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                                                                                                                                                                                                                                                       03-OCT-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       09-AUG-2001
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABK39743 standard; CDNA; 2442 BP.
CDNA encoding clone #48005 (L979P) of lung tumour protein version
20020202514-A2.
17-JAN-2002.
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                       Score 28.2; DB 11; Length 1799;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                 Score 28.2; DB 3; Length 1863;
Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                            (SCRI ) SCRIPPS RES INST.
(SYGN ) SYNGENTA PARTICIPATIONS AG.
ry Match
t Local Similarity 61.6%; Pred. No. 2.88+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  23.SEP-1999.
(RADO/) RADOSEVICH J A. ery Match 10.3%; Score 28.2; DB 2; Length 2442; ery Match similarity 54.3%; Pred. No. 3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 2442;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%; Score 28.2; DB 3; Length 2250; 54.3%; Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 2442
                                                                                                Length 1465
                                                                                                                                                                                                                                                                                                                                                                               ABZ16408 standard; DNA; 2000 BP.
Arabidopsis thaliana stress regulated gene SEQ ID NO 4213.
WO200216655-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human lung cancer associated cDNA L979P extended sequence US2002197669-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ACA03258 standard; cDNA; 2442 BP.
Lung cancer therapyand diagnosis associated cDNA #1745.
US2002172952-A1.
                                                                                                                                                                                                                                                              AAC41185 standard; DNA; 1863 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 38339.
BP1033405-A2.
06-SEP-2000.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAC49336 standard; DNA; 2250 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 60775
EP1033405-A2.
                AAS68406 standard; cDNA; 1465 BP.
DNA encoding novel human diagnostic protein #4210.
W0200175067-A2.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%; Score 28.2; DB 8; 54.3%; Pred. No. 3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.3%; Score 28.2; DB 6; 54.3%; Pred. No. 3e+02;
                                                                                             10.3%; Score 28.2; DB 5; 59.3%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAZ23609 standard; cDNA; 2442 BP.
Human labyrinthin cDNA.
WO9947683-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ACA12072 standard; cDNA; 2442 BP
                                                                                                                                              ADI30894 standard; cDNA; 1799 BP
                                                                                                                                                                                                                          10.3%;
51.2%;
                                                                                                                                                                                                                                                                                                                                   10.3%;
54.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          26-DEC-2002.
(BANG/) BANGUR C S.
(FANG) FANGER G R.
(WANG/) WANG A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (SWIT/) SWITZER A P. (MCNE/) MCNEILL P D. (CLAP/) CLAPPER J D.
                                                                                                                                                                                           19-AUG-2003.
(INCY-) INCYTE CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 914
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 911
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RESULT 913
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
RESULT 910
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Best Local Similarity
RESULT 908
                                                                                                            Best Local Similarity RESULT 907
                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
RESULT 909
                                                                                                                                                           Human cDNA #220.
US6607879-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          06-SEP-2000.
                                                                                                                                                                                                                                                                                                                                                                                                                                28-FEB-2002.
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(WANG/) V
(WANG/) V
(SWIT/) 8
                                                                                                Query Match
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      RESULT 906
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DE
PN
PA
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ABL10930 standard; cDNA; 3365 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 27272.
WO200171042-A2.
                                                                                                                                                  Length 2442;
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 3038;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 3038;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 3497;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 3145;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 3365;
10.3%; Score 28.2; DB 8; Length 2442; 54.3%; Pred. No. 3e+02;
                                                                                                                                                                                                                                                                                                    Length 2949;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 3218,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           cDNA encoding a protein similar to human sorting nexin.
WO200262839-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABAUB779 standard; cDNA; 3497 BP.
Human RING-H2 homologue-encoding cDNA, SEQ ID NO:555.
WOZOUS7188-A2.
                                                                                                                                                                                                                                                                                                                                                       AD182490 standard; DNA; 3038 BP.
Human modifier of p21 (MP21) gene sequence SeqID56
WO2004005486-A2.
                                                                                                                                                    10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 28.2; DB 12;
Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 28.2; DB 12;
Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                  Score 28.2; DB 8;
Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 28.2; DB 4;
Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 28.2; DB 4;
Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         28.2; DB 6;
No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28.2; DB 2;
Pred. No. 3.3e+02;
                                                                                                                                                  Score 28.2; DB
Pred. No. 3e+02;
                                                      ADH47310 standard; cDNA; 2442 BP.
Human lung tumour cDNA clone, SEQ ID No 1791.
08-MAY-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADD29717 standard; mRNA; 3507 BP.
Mouse tumour suppressor mRNA SEQ ID NO:173.
WO2003058201-A2.
17-J7U-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score
Pred.
                                                                                                                                                                                                      ACAS2736 standard; DNA; 2949 BP.
Prokaryotic essential gene #34393
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Antipsoriatic cDNA sequence #607.08-APR-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CDNA; 3218 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABQ79518 standard; cDNA; 3145 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WO2004020.
08-APR-2004.
(GETH ) GENENTECH INC.
10.3%;
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(EDGE/) EDGERTON M D.
                       (CHOM/) CHOMET P S. (LACC/) LACCETTI L B.
                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                          Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADM47730 standard; DNA; 7858 BP.
Polynuclectide sequence #148 useful in producing transgenic plants.
US2003233670-A1.
(QUAR-) QUARK BIOTECH INC.
(CLEV-) CLEVELAND CLINIC FOUND.
rry Match
st Local Similarity 51.2%; Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                               ADA19288 standard; cDNA; 3598 BP.
Human insulin resistance marker IRM148 (ring finger protein 3)
WO200298355-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 28.2; DB 10; Length 3921; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.3%; Score 28.2; DB 10; Length 3598; 51.2%; Pred. No. 3.4e+02;
                                                                                                                                                            Length 3527;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.3%; Score 28.2; DB 6; Length 6351; 52.1%; Pred. No. 4.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                      Length 3527;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 5563;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 7065;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABL33746 standard; DNA; 6351 BP.
Human immune system associated gene SEQ ID NO: 1719.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL33553 standard; DNA; 7065 BP.
Human immune system associated gene SEQ ID NO: 1526
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.3%; Score 28.2; DB 2; 64.6%; Pred. No. 4e+02;
                                                                                                                                                         Score 28.2; DB 2;
Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                      Score 28.2; DB 6;
Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 28.2; DB 6;
Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Enterococcus faecalis polynucleotide #3265.
US6617156-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (MEDI-) MEDICAL COLLEGE GEORGIA RES INST.
                                                                             AAV61482 standard; cDNA; 3527 BP.
Human secreted protein cw775_1 cDNA.
WO9841539-A2.
                                                                                                                                                                                                ABG92009 standard; cDNA; 3527 BP.
Human polynucleotide SEQ ID NO 6.
US2002065394.Al.
30-MAY-2002.
(JACO/) JACOBS K.
(MCCO/) MCCOY J. M.
(LAVA/) LAVALLIE E R.
(COLL/) COLLINE R.C.
(EVAN/) EVANS C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAX08941 standard; DNA; 5563 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADH85380 standard; DNA; 3921 BP
                                                                                                                                                         10.3%;
51.2%;
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51.2%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                09-SEP-2003.
(DOUC/) DOUCETTE-STAMM L A.
(BUSH/) BUSH D.
                                                                                                                         24-SEP-1998. (GEMY ) GENETICS INST INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        GATA-1 promoter region.
WO9856902-A2.
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        12-DEC-2002.
(CLIN-) CLINGENIX INC.
                                                                                                                                                                                                                                                                                                                                        (MERB/) MERBERG D.
(TREA/) TREACY M.
(SPAU/) SPAULDING V.
                                             Best Local Similarity RESULT 924
                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 926
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
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RESULT 930
                                                                                                                                                                     Local Similarity
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                                  Query Match
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RESULT 925
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Human immune/haematopoietic antigen genomic sequence SEQ ID NO:27748.
W0200157182-A2.
09-A40G-2001.
(HUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28.2; DB 4; Length 14169;
st Local Similarity 46.6%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                   ABL13258 standard; cDNA; 9936 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 34256.
WC200171042-A2.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                            Human chemically modified disease associated gene SEQ ID NO 242 WO200200927-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             03-JAN-2002.
(EDIG-) EPIGENOMICS AG.
ery Match 10.3%; Score 28.2; DB 6; Length 13503;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human immune system associated gene SEQ ID NO: 191.
W020020928-A2.
03-JAN-2002.
(BPIG-) EPIGENOMICS AG.
10.3%; Score 28.2; DB 6; Length 11422; st Local Similarity 49.7%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 11863;
             Length 7858;
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                                                                                                                                                                                          Length 8067;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human chemically pretreated gene sequence #72 strand W0200202806-A2.
10-JAN-2002.
[BPIG:) EPIGENOMICS AG.
10.3%; Score 28.2; DB 6; Lenget Local Similarity 47.9%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                           ABKX9936 standard; DNA; 11422 BP.

Human chemically pretreated gene sequence #9 strand
W0200202806-A2.

10-JAN-2002.

(EPIG-) EPIGENOMICS AG.

10-JA Match

10.3%; Score 28.2; DB 6; Len
ist Local Similarity 49.7%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABL34070 standard; DNA; 13503 BP.
Human immune system associated gene SEQ ID NO: 2043
WO200200928-A2.
Query Match
Best Local Similarity 54.3%; Pred. No. 4.5e+02;
RESULT 932
                                                                                                                                   mrarolog.
(EPTG-) EPIGENOMICS AG.
(EPTG-) EPIGENOMICS AG.
lery Match 10.3%; Score 28.2; DB 6;
                                                                                                                                                                                                                                                                                                                                                                  10.3%; Score 28.2; DB 4; 61.6%; Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.2; DB 8;
Pred. No. 5.3e+02;
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Human heparin cofactor II (HCF2) gDNA.
WO2003091453-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL32218 standard; DNA; 11422 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DNA; 11863 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABK40062 standard; DNA; 13784 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAK72936 standard; DNA; 14169 BP.
                                                                              ABN80225 standard; DNA; 8067 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                C. elegans ses-3 DNA.
WO2003000717-A2.
                                                                                                                                                                                                        Best Local Similarity
RESULT 933
                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 934
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 935
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABX94628 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            03-JAN-2003.
(ELEG-) ELEGENE AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 937
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 938
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Length 96599;

Length 96599;

Length 96599;

Length 110000;

Length 110000;

Length 110000;

Length 110000

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Score 28.2; DB 10; Length 137870;
Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ACAIBIGO standard, DNA, 247 BP.
Prokaryotic essential gene antisense oligonucleotide #6030.
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AASS0893 standard; DNA; 247 BP.
Staphylococcus aureus cellular proliferation inhibitory
WO200170955-A2.
                                                                                                                                                                                                                                                                                                                                      ADM74576 standard; DNA; 96599 BP.
Murine carcinoma associated (CA) nucleic acid #124
US2004072154-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       FRAILS.
27-0000.
(CNRS ) CNRS CENT NAT RECH SCI.
(IFRE-) IFREMER INST FR RECH EXPL MER.
10.3%; Score 28.2; DB 5;
Lery Match 10.3%; Score 28.2; DB 5;
                                                                                                                                  10.3%; Score 28.2; DB 10; 51.2%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                  Score 28.2; DB 10;
Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Z7-CC1-z0CC.
(CNRS.) CNRS CENT NAT RECH SCI.
(IFRE-) IFREMER INST FR RECH EXPL MER.
10.3%; SCOTE 28.2; DB 5;
str Match
10.3%; Pred. No. 1.7e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 28.2; DB 12;
Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             28.2; DB 2;
No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        28.2, DB 6;
No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 28.2; DB 2;
Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 28.2; DB 2;
Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%; Score 28; DB 4; 50.8%; Pred. No. 1.5e+02;
     51.2%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 28.2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           #3.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAH41225 standard; DNA; 349980 BP.
Pyrococcus abyssi genomic fragment #4.
FR2792651-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score Pred. 1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAH41224 standard, DNA; 349980 BP. Pyrococcus abyssi genomic fragment FR2792651-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Pred.
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Human matrilin-3 gene SEQ ID NO:1.
Query Match
Best Local Similarity 59.3%; Preor
                                                                                                                                                                                    ADC85461 standard; DNA; 96599 BP.
Mouse Map2k5 genomic sequence.
WO2003045230-A2.
                                          ADB72719 standard; DNA; 96599 BP
                                                                                                                                                                                                                                      MULACY-
05-JUN-2003.
(SAGR-) SAGRES DISCOVERY.
ALV MATCh 10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.3%;
54.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.3%;
53.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.3%;
50.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%;
51.2%;
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                                                                                              30-JAN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                           15-APR-2004.
(MORR/) MORRIS D W.
(ENGE/) ENGELHARD E K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Local Similarity
Best Local Similarity
RESULT 950
                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 952
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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                                                       Mouse Map2k5 gene.
WO2003008583-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              27-OCT-2000
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                                                                                                                                    Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 24001;
                                          Length 15657;
                                                                                                                                                                                                                                                                                                                                           Length 15849;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 20860;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 80073;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 18887;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 20860;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 9; Length 96599;
                                                                                                                                                                                             Length 15848
                                                                                            AAZ32190 standard; DNA; 15848 BP.
Human heparin cofactor II exon 1 to 5 nucleotide sequence
WO9950454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             SEQ ID NO:1499
 06-NOV-2003.
(GENA-) GENAISSANCE PHARM INC.
ery Match
ery Match
(GENA-) cimilarity 45.9%; Pred. No. 5.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAX13292 standard; DNA; 20860 BP.
Enterococcus faecalis genome contig SEQ ID NO:355
WO9850555-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 28.2; DB 12;
Pred. No. 6.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%; Score 28.2; DB 10; 47.9%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADF11646 standard; DNA; 94752 BP.

Human chromosome 17 clone HPRC905N1 nucleic acid
WO2003087763-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             MUNDSOLOUS
12-NOV-1998.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
(ELY MATCH 10.3%; Score 28.2; DB 2;
(ELY MATCH 10.18) SO.4%; Pred. No. 6.5e+02;
                                                                                                                                                       07-OCT-1999.
(WHED ) WHITEHEAD INST BIOMEDICAL RES.
ery Match 10.3%; Score 28.2; DB 2;
ery Match 45.9%; Pred. No. 5.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 28.2; DB 6;
Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 28.2; DB 2;
Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                 11.APR-2002.

11.APR-2002.

(GENE-) GENE LOGIC INC.

ery Match

10.3%; Score 28.2; DB 6;

ery Match

15.9%; Pred. No. 5.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                               WCCCT-2001.
(GENA-) GENAISSANCE PHARM INC.
(GENA-) GENAISSANCE 10.3%; Score 28.2; DB 6;
ery Match 10.3%; Pred. No. 6.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                             AASI6543 standard; DNA; 18887 BP.
DNA encoding UDP glycosyltransferase 1 (UGT1A1)
WO200179230-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AD022771 standard; DNA; 24001 BP.
DNA encoding human interleukin 22 receptor.
US2004097447-A1.
2.0-MAY-2004.
(ISIS-) ISIS PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABS99087 standard; DNA; 20860 BP.
Enterococcus faecalis contig sequence #355.
US2002120116-Al.
29-AUG-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.3%; Score 28.2;
                                                                                                                                                                                                                                            ABN95864 standard; DNA; 15849 BP.
Gene #2362 used to diagnose liver cancer.
WO200229103-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Mouse Map2ks carcinoma associated gene, WO2003057146-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADA02981 standard; DNA; 96599 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           23-OCT-2003.
(CELL-) CELLTECH R & D INC.
(UYRO-) UNIV ROTTERDAM RASMUS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%;
61.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              17-JUL-2003.
(SAGR-) SAGRES DISCOVERY.
                                            Ouery Match
Best Local Similarity
RESULT 941
                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 943
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 945
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (KUNS/) KUNSCH C A.
(DILL/) DILLON P J.
(BARA/) BARASH S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 946
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 948
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 949
                                                                                                                                                                                                             Best Local Similarity RESULT 942
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 944
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
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                                                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                               Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
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sequence #2117.

Length 349980;

Length 349980;

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27-SEP-2001.
(ELIT-) ELITRA PHARM INC.
   WO200170955-A2.
                                                                                                                                                                27-SEP-200
                                                                                                                                                                                                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
                                                          Query Match
                                                                                                                                                                                                    Query Match
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                                                                                                                                                                                                                                      RESULT 972
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AASS5091 standard; DNA, 441 BP.
Staphylococcus aureus DNA for cellular proliferation protein #1403.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAS51903 standard; DNA; 438 BP.
Staphylococcus aureus DNA for cellular proliferation protein #320.
WO200170955-A2.
                                                                                                                                                                                                                          ARK91741 standard; DNA; 327 BP.

DNA encoding murine glucose transport-related polypeptide
WO200233046-A2.
25-APR-2002.
(UTMA-) UNIV MASSACHUSETTS.
ery Match
st Local Similarity 50.8%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 TACACACOLO.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

ery Match

10.3%; Score 28; DB 5; Length 418;
                                   Length 247;
                                                                                                                                                                               Length 268,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 8; Length 369;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 387;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 373;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.3%; Score 28; DB 4; Length 438; 50.8%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 438;
                                                                                   ABLB6380 standard; cDNA; 268 BP.

Human ovarian cancer related cDNA clone SEQ ID NO:9358.
WC200192581-A2.
(G-DEC-2001.
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                Human gene trapped sequence (GTS) cDNA SEQ ID NO 784, US2002110809-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Human ovarian antigen HPDVL36 cDNA, SEQ ID NO:1913 WO200200677-A1.
       (ELIT-LECT.)

SELY MATCH

10.3%; Score 28; DB 8; I

try Match

tr Local Similarity 50.8%; Pred. No. 1.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAC28678 standard; cDNA; 373 BP.
Human secreted protein 5' EST, SEQ ID NO: 32753.
EP1033401-A2.
06-SEP-2000.
                                                                                                                                                                               Score 28; DB 6;
Pred. No. 1.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%; Score 28; DB 3; 166.7%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.3%; Score 28; DB 6; 58.2%; Pred. No. 1.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.3%; Score 28; DB 8;
53.2%; Pred. No. 1.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%; Score 28; DB 8; 50.8%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABV16971 standard; cDNA; 418 BP.
Human prostate expression marker cDNA 16962.
WC200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ACF75022 standard; DNA; 438 BP.
Staphylococcus aureus DNA #2702.
WO200294868-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABQ56033 standard; cDNA; 387 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             03-JAN-2002.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                               10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   27-SEP-2001.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                   (NEHL/) NEHLS M C.
(ZAMB/) ZAMBROWICZ B.
(SAND/) SANDS A T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
BESUL 967 1 EAVIEGT CD
DE Human prostate expres
PD 23-AUG-2001.
PA (MILL-) MILLENNIUM PR
                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 964
ID ABK5117 standard; cl
DE HUMAN Gene trapped 6:
PN US2002110809-A1.
PD 15-AUG-2002.
PA (ZAMB/) NEHLS M C.
PA (ZAMB/) SANDS A T.
                                             Best Local Similarity RESULT 962
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 969
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 970
                                                                                                                                                                                            Best Local Similarity
RESULT 963
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 965
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity RESULT 966
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 968
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  28-NOV-2002.
(CHIR-) CHIRON SPA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (GEST ) GENSET.
03-OCT-2002
                                                                                                                                                                                                                                                                                                                           Query Match
                                 Query Match
                                                                                                                                                                               Query Match
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AASS5314 standard; DNA; 441 BP.
Staphylococcus aureus DNA for cellular proliferation protein #1626.
WO200170955-A2.
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ABQ33825 standard; DNA; 573 BP.
Oligonucleotide for detecting cytosine methylation SEQ ID NO 20416.
WO200218632-A2.
O7-WAR-2002.
(FPIG-) EPIGENOMICS AG.
10.3%; Score 28; DB 6; Length 573;
erry Match
10.3%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABQ33824 standard; DNA; 573 BP.
Oligonucleotide for detecting cytosine methylation SEQ ID NO 20415.
WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    A ABQ35515 standard; DNA; 573 BP.

3 Oligonucleotide for detecting cytosine methylation SEQ ID NO 22106.

4 NO20021632-A2.

5 O7-MAR-2002

A (EPG-) EPIGENOMICS AG.

Query Match

10.3%; Score 28; DB 6; Length 573;

Best Local Similarity 52.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABQ35514 standard; DNA; 573 BP.
Oligonucleotide for detecting cytosine methylation SEQ ID NO 22105.
WO200218632-A2.
                                                                                                                                                                                                                                                       AASS4490 standard, DNA, 441 BP.
Staphylococcus aureus DNA for cellular proliferation protein #802.
WO20170955-A2.
27-SEP-2001.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                             10.3%; Score 28; DB 4; Length 441; 50.8%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 441;
    Length 441;
                                                                                                                                                                                           Length 441,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 573
10.3%; Score 28; DB 4; 50.8%; Pred. No. 1.9e+02;
                                                                                                                                                                                      10.3%; Score 28; DB 4; 50.8%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 28; DB 8;
Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.3%; Score 28; DB 5;
r ronal Similarity 50.0%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.3%; Score 28; DB 5;
or Tonal Similarity 50.0%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.3%; Score 28; DB 6; 52.6%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human prostate expression marker cDNA 45421. 93.8 Human prostate expression marker cDNA 45421.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human prostate expression marker cDNA 36399 WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Prokaryotic essential gene #1679.
020027183-A2.
03-0CT-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABV36408 standard; cDNA; 528 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA; 441
                                                                                                                                                                (ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            07-MAR-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                             Local Similarity
              Best Local Similarity RESULT 971
                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 973
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 974
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Best Local Similarity
RESULT 975
                                                                                                                                                                                                                                                                                                                                                                                                                                                  ACA20022 standard;
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Query Match

Query Match

DE BA

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ADO43421 standard; cDNA; 826 BP.
Lutzomyia longipalpis (sand fly) salivary protein LJM114 cDNA
WO2004039958-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 826;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.3%; Score 28; DB 8; Length 1002; 50.0%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 783;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             23-5405-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 10.3%; Score 28; DB 5; Length 764;
                                                                                                        Length 756;
                                                                                                                                                                                                                                                                                                                                Length 764;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human breast cancer expressed polynucleotide 16294 WO200151628-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DNA encoding Acinetobacter baumannii protein #870 US6562958-B1.
WO200151628-A2.
19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
10.3%; Score 28; DB 4; I erv Match
                                                                                                                                                                                                                                                                      23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.3%; Score 28; DB 5; I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Klebsiella pneumoniae polynucleotide seqid 1532.
US6610836-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PD 13-MAY-2003.

PA (GENO-) GENOME THERAPEUTICS CORP.

QUERY MAtch

10.3%; Score 28; DB 9; 1

Best Local Similarity 52.6%; Pred. No. 2.3e+02;

RESULT 993
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.3%; Score 28; DB 5; 1 50.0%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     26-AUG-2003.
(GENO-) GENOME THERAPEUTICS CORP.
(ery Match 10.3%; Score 28; DB 11;
ery Match 56.5%; Pred. No. 2.38+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                19-JUL-2001.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

ery Match

10.3%; Score 28; DB 4;

or rocal Similarity 47.4%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.3%; Score 28; DB 12; 58.3%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABV15622 standard; cDNA; 895 BP.
Human prostate expression marker cDNA 15613.
WO200160860-A2,
                                                                                                                                                                                                                                                                                                                                                                                                            ABV24119 standard; cDNA; 764 BP.
Human prostate expression marker cDNA 24110
WO200160860-A2.
                                                                                                                                                                                                              Human prostate expression marker cDNA 29989
WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        13-MAY-2004.
(USSH ) US DEPT HEALTH & HUMAN SERVICES.
(PESQ-) CENT PESQUISAS GONCALO MONIZ.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Prokaryotic essential gene #26382.
WO200271183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADF02733 standard; DNA; 1062 BP.
Bacterial polynucleotide #3018.
                                                                                                                                                                                             ABV29998 standard; cDNA; 764 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       CDNA; 788 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA; 819 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DNA; 783
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (ELIT-) ELITRA PHARM INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
RESULT 994
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity
RESULT 991
                                                                                                                             Best Local Similarity RESULT 988
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAL23837 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADA29583 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ACH95737 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                03-OCT-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                              Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 995
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    RESULT 992
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         RESULT 996
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABK63310 standard; cDNA; 603 BP. Rat sequence differentially expressed in response to a hepatotoxin #1217. WO200210453-A2. 07-FEB-2002. (GENE-) GENE LOGIC INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               NO:2668.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Primary rat hepatocyte toxicity modelling related gene SEQ ID WO2003065993-A2.
                                                                                                                                                          ABQ57311 standard; cDNA; 582 BP.
Human colon cancer related nucleotide sequence SEQ ID NO:1006.
WO200229086-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 599;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 603
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 603
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.3%; Score 28; DB 4; Length 665; 77.3%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 665;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.3%; Score 28; DB 6; Length 603; 52.6%; Pred. No. 2.1e+02;
                                                                                    Length 573;
                                                                                                                                                                                                                                                                                                            Length 582;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAL22994 standard, cDNA, 756 BP.
Human breast cancer expressed polynucleotide 15451
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.3%; Score 28; DB 5; 1
77.3%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WOZOCCO.
07-MUG-2003.
(GENE ) GENE LOGIC INC.
(GENY Match
10.3%; Score 28; DB 10;
10.3%; Score 28; DB 10;
10.3%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                               10.3%; Score 28; DB 6; 59.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.3%; Score 28; DB 12; 50.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.3%; Score 28; DB 10; 52.6%; Pred. No. 2.1e+02;
                                                                                       Score 28; DB 6;
Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                             ACH69966 standard; DNA; 599 BP.

Human genome derived single exon probe #3161.
US2003194704-A1.
16-OCT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-10-0CT-20019-0CT-20019-0CT-20019-0CT-20019-0CT-20019-0CT-20019-0CT-20019-0CT-2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Porcine DNA encoding interleukin-18 (IL-18)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       26-JUN-2001.
(NORD, ) NORINGUISANSHO KACHIKU EISEI.
(MUNE/) WINDELA Y.
(MORI/) MORI Y.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (NORQ ) NORINSUISANSHO KACHIKU EISEI
(WINE) MUNETA Y.
(MORI/) MORI Y.
(SHIM/) SHIMOCHI Y.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADB57606 standard; DNA; 603 BP.
Toxicity-related gene, SEQ ID 2632.
WO2003064624-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADB52126 standard; DNA; 603 BP
                                                                                       10.3%;
52.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AA166519 standard; DNA; 665
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      14-AUG-2003.
(GENE-) GENE LOGIC INC.
                                  07-MAR-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                        Best Local Similarity
RESULT 980
ID ARQ57311 standard; CL
DE Human colon cancer re
PN WC200229086-A2.
PD 11-ARR-2002.
PA (FARB ) BAYER CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Pig coding sequence. JP2001169785-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
RESULT 983
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 986
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (MORI/) MORI Y.
(SHIM/) SHIMOCHI Y.
(ARAI/) ARAI K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 982
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
RESULT 984
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 985
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                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 981
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ARAI K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    17-APR-2001.
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Query Match

Match

Ouery

(MORI/) (SHIM/) (ARAI/)

Length 2563;

Length 2841;

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LUCLY MATCH 10.3%; Score 28; DB 12; Length 3224; Best Local Similarity 50.8%; Pred. No. 3.8e+02; RESULT 1012
                                                                                                                                                                                                                                                                                                                                     Score 28; DB 6; Length 2871;
Pred. No. 3.7e+02;
                                                                                                                                                                                                                                            ABV94250 standard; CDNA; 2871 BP.
Breast carcinoma related nucleotide sequence SEQ ID NO:241.
9200246467-A2.
13-UNA-2002.
(IPSO-) IPSOGEN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AD019407 standard; DNA; 3127 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 2226.
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADQ23637 standard; DNA; 3224 BP.
Human soft Lissue sarcoma-upregulated DNA - SEQ ID 6457
WO2004048938-A2.
10-JUN-2004.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity ......
IREGULT 1009
ID Hombospondin 3 (THBS3) mRNA, complete cds DNA
PN WO2003072827-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAX83478 standard; cDNA to mRNA; 3960 BP.
A.thaliana potassium channel KAT2 gene.
W099125-A1.
24-UUN-1999.
(PLAC ) MAX PLANCK GES FOERDERUNG WISSENSCHAFTEN.
                                                         10.3%; Score 28; DB 12; 50.8%; Pred. No. 3.5e+02;
                                                                                                                                                                                               Score 28; DB 8;
Pred. No. 3.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                  04-SEP-2003.

(CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT.

ery Match 10.3%; Score 20; DB 11;

11 Innal Similarity 50.8%; Pred. No. 3.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        04-SEP-2003.
(CHIL-) CHILDREN'S HOSPITAL MEDICAL CENT.
ery Match
10.3%; Score 28; DB 11;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10-UU-2004.

(PROT-) PROTEIN DESIGN LABS INC.

ery Match 10.3%; Score 28; DB 12;

ery Match 50.8%; Pred. No. 3.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.3%; Score 28; DB 2; ] 55.0%; Pred. No. 4.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  WOZUULLOZOU.
08-MAR_2001.
(SUNTORY LITD.
(SAOK ) NIPPON PAPER IND CO LID.
(SAOK ) "IPPON PAPER IN CO LID.

"Match "logic Self"; Bred. No. 4e+02;
"Totty 52.6%; Pred. No. 4e+02;
                                                                                                                                                                                                                                                                                                                                                                       Aurosovo standard; DNA; 2871 BP.
Human thrombospondin 3 (THBS3) DNA sequence.
WO2003072827-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Histdine protein kinase coding sequence #3.
WO200116332-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAH14686 standard; cDNA; 4281 BP.
Human cDNA sequence SEQ ID NO:12384.
                                                                                                                          Prokaryotic essential gene #12052.
WO200277183-A2.
                                       (REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                          DNA; 2841 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           A (PROT-) PROTEIN DESIGN LABS INC.
Query Match 10.3%; Sco
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAF75866 standard; DNA; 3708 BP.
                                                                                                                                                                                               10.3%;
51.6%;
                                                                                                                                                                                                                                                                                                                                     10.3%;
                                                                                                                                                                           (ELIT-) ELITRA PHARM INC.
                                                                      Best Local Similarity RESULT 1006
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 1010
                                                                                                                                                                                                             Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
                                                                                                          ACA30395 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1011
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 1013
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 1014
     EP1396543-A2.
                                                                                                                                                             03-OCT-2002
                                                       Query Match
                                                                                                                                                                                                                                                                                                                                     Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                               Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                 RESULT 1007
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    P P R E P
                                                    Length 1062;
                                                                                                                                                                                           DB 5; Length 1431;
                                                                                                                                                                                                                                                                                                                             10.3%; Score 28; DB 5; Length 1431; 56.5%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 1431;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.3%; Score 28; DB 5; Length 1431; 56.5%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 2304;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 2114;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.3%; Score 28; DB 3; Length 2158; 49.3%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%; Score 28; DB 4; Length 2563; 50.8%; Pred. No. 3.5e+02;
                                                                           DNA encoding novel human diagnostic protein #18175.
MO200175067-A2.
11-OCT-2001.
iry Match
                                                                                                                                                                                                                          DNA encoding novel human diagnostic protein #30043
                                                                                                                                                                                                                                                                                                                                                                                AAS82055 standard; cDNA; 1431 BP.
DNA encoding novel human diagnostic protein #17859.
WO200175067-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         cDNA; 1431 BP. human diagnostic protein #28838
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AACS9077 standard; cDNA; 2114 BP.
Human secreted protein coding sequence SEQ ID NO:
WC200055171-A1.
          12-AUG-2003.
(GENO-) GENOME THERAPEUTICS CORP.
ery Match
10.3%; Score 28; DB 10;
''^~al Similarity 50.0%; Pred. No. 2.5e+02;
                                                                                                                                                                                         10.3%; Score 28; DB 5; 56.5%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.3%; Score 28; DB 3; 1 49.3%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 28; DB 8; I
Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%; Score 28; DB 5; 56.5%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAC93380 standard; cDNA; 2158 BP.
Human secreted protein gene 17 SEQ ID NO:27.
WO200061620-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAK94255 standard; cDNA; 2563 BP.
Human full-length cDNA, SEQ ID NO: 2868.
EP1130094-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADL30835 standard; cDNA; 2563 BP.
Full length human cDNA clone SeqID 2868.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Prokaryotic essential gene #34480.
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ACA52823 standard; DNA; 2304 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                21-SEP-2000.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  19-OCT-2000.
(HUMA-) HUMAN GENOME SCI INC.
(ROSE/) ROSEN C A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  05-SEP-2001.
(HELI-) HELIX RES INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 1001
DE AAC59077 standard; cf
DE Human secreted protein
PD VC200055171-A1.
PD 21-SEP-2000.
PA (HUMA-) HUMAN GENOME
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAS93034 standard; c
DNA encoding novel h
WO200175067-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
RESULT 1004
                                                         Best Local Similarity
RESULT 997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
RESULT 1002
                                                                                                                                                                                                      Best Local Similarity RESULT 998
                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 999
                                                                                                                                                                                                                                                                                         11-OCT-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (HYSE-) HYSEQ INC.
Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                 11-OCT-2001.
(HYSE-) HYSEQ INC.
US6605709-B1.
                                                  Query Match
                                                                                                                                                                                         Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
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                                                                                                                                                                                                                                            DE NO.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DE DE LO
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ID PN PD PA

Length 2871;

Length 3127;

Length 3127;

Length 3708;

Length 3960;

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Genomic sequence #541 encoding for novel human respiratory antigen. 02-AUG-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 1030
ID ART42839 standard; DNA; 9137 BP.
DE Human nucleic acid-associated protein (NAAP) coding sequence #20.
PN W02003010329-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADG41897 standard; DNA; 7461 BP.

Human respiratory system associated genomic DNA seq id 1135.

VG200315893-A1.

20-NOV-2003.

(HUMA-) HUMAN GENOME SCI INC.

Query Match

Best Local Similarity 50.0%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.3%; Score 28; DB 10; Length 8602; 63.2%; Pred. No. 5.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABA21227 standard, DNA, 9042 BP.
Human nervous system related polynucleotide SEQ ID NO 13558.
WO20159063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                            Length 7461;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 7461;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 9042;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 9415;
                                                  Length 7461;
                                                                                                                                                                                                              Length 7461;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (HORT-) HORTICULTURE & FOOD RES INST NEW ZEALAND (STAN/) STANLEY D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          WOZUWJZZZOJ.
06-FEB-2003.
(INCY-) INCYTE GENOMICS INC.
(INCY-) INCYTE GENOMICS INC.
10.3%; Score 28; DB 8; L
HETY MATCH 55.0%; Pred. No. 5.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 5; I
lery Match 10.3%; Score 28; DB 5; I
                                                    10.3%; Score 28; DB 4; I 50.0%; Pred. No. 5.2e+02;
                                                                                                                                                                                       (HUMA-) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 50.0%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          USACCO.

13-MAR-2003.

(HUMA-) HUMAN GENOME SCI INC.

(HUMA-) HUMAN GENOME SCI INC.

10.3%; Score 28; DB 9;

cery Match 50.0%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.3%; Score 28; DB 6; 50.8%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                              10.3%; Score 28; DB 8; 50.0%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                          RESULT 1025

RESULT 1025

ID ACADAJOCA standard; DNA; 7461 BP.

DD DNA encoding human lung cancer antigen HIPAA05.

PN US2002173454-A1.

PD 21-NOV-2002.

PA (ROBE/) ROSEN C A.

PA (RUBE/) RUBEN S M.

PA (RUBE/) RARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADB96750 standard; DNA; 7461 BP.
Novel lung cancer antigen genomic DNA #21.
US2003049703-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADC36595 standard; DNA; 8602 BP. DNA sequence of the invention #11 WO2003027141-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        BP
                                                                                                              AAS28701 standard; DNA; 7461 BP.
              02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DNA; 9415
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Ouery Match
Best Local Similarity
RESULT 1032
                                                                    Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 1027
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 1031
ID ABL32294 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                          Best_Local Similarity
RESULT 1026
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             03-APR-2003
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                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human immune/haematopoietic antigen genomic sequence SEQ ID NO:40745.
WO200157182-A2.
                                                                                                                                                                                     ABL26736 standard; DNA; 6883 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 31681
WO200171042-A2.
27-SEP-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 5886;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.3%; Score 28; DB 6; Length 4769; 55.0%; Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MILL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.3%; Score 28; DB 5; Length 4911;
                                             Query Match 10.3%; Score 28; DB 4; Length 4281; Best Local Similarity 55.0%; Pred. No. 4.2e+02; RESULT 1015
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match imilarity 63.2%; Pred. No. 4.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 6883;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 6176;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAS30639 standard; DNA; 7461 BP.
DNA encoding novel lung cancer antigen, Seq ID No 91.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Zea mays alanine amino transferase (aat1) intron 9 WO200175071-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABQ60834 standard; cDNA; 4769 BP.
FLJ10898 fis clone NT2RPS003492 encoding sequence.
0.00231111-A2.
18-APR-2002.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADP90748 standard; DNA; 5886 BP.
Human hepatic-fibrosis disease marker SEQ ID 210.
JP200259877-A.
16-SEP-2003.
(SUMU) SUMITOMO SEIYAKU KK.
(SUMU) SUMITOMO 51038; Score 28; DB 10; I St.
Local Similarity 55.08; Pred. No. 4.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         MOLOCIO
11-00-12001.
(REGC ) UNIV CALIFORNIA.
(REGC ) UNIV CALIFORNIA.
(REGC ) UNIV CALIFORNIA.
(REGC ) UNIV 47.2%; Pred. No. 4.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.3%; Score 28; DB 4; 53.7%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Jr 1018
ADL460708 standard; DNA; 4911 BP.
Human ovarian cancer DNA marker #19968.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity 55.0%; Pre RESULT 1021
ID AAD21947 standard; DNA; 6176 BP.
                                                                                                              ADJ75040 standard; DNA; 4281 BP.
Marker gene SEQ ID NO:292.
EP1394274-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAK85933 standard; DNA; 5742 BP
                07-FEB-2001.
(HELI-) HELIX RES INST
                                                                                                                                                                          (GENO-) GENOX RES INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
RESULT 1023
ID AAS30639 standard,
DE DNA encoding novel
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
RESULT 1020
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RESULT 1022
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EP1074617-A2.
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Query Match

Query Match

Query Match

Query Match

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ADF74520 standard, DNA; 28320 BP.
Human genomic DNA encoding the cholesteryl ester transfer protein (CETP).
WO2003091698-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAL36280 standard; DNA; 32174 BP.
Human musculoskeletal system related polynucleotide SEQ ID NO 2645.
WO200155367-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AALO7447 standard; DNA; 32174 BP.

Human reproductive system related antigen DNA SEQ ID NO: 10135.

MO200155320-A2.

O2-AUG-2001.

A (HUMA-) HUMAN GENOME SCI INC.

Query Match

Best Local Similarity 50.0%; Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             BEBL LOLG.

BESULT 1048

ID AAL03792 standard; DNA; 32174 BP.

DE Human reproductive system related antigen DNA SEQ ID NO: 6480.

BD 02-AUG-2001.

PD 02-AUG-2001.

PA (HUMA-) HUMAN GENOME SCI INC.

PA (HUMA-) HUMAN GENOME SCI INC.

Ouery Match

Ouery Match

IO.3*; Score 28; DB 4; Length 32174;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human reproductive system related antigen DNA SEQ ID NO: 10179 WO200155320-A2.
                                                                                                                                                                                                               ADG98369 standard; DNA; 28313 BP.
Human cholesteryl ester transfer protein (CETP) gene.
lery Match
10.3*; Score 28; DB 10; Length 28313;
sst Local Similarity 60.5*; Pred. No. 8.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                Length 28320;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match
.ery Match
.ery Match 10.3%; Score 28; DB 4; Length 32174;
.ery Match 50.0%; Pred. No. 8.7e+02;
        Length 23683;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 32174;
                                                                                                                                                              Length 23683;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 32174;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     RESULT 1045
ID AAI62606 standard; DNA; 32174 BP.
DE Human breast or ovarian antigen genomic DNA SEQ ID NO:
PN WO200155324-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              90
                                                              DNA; 23683 BP.
cell signalling DNA sequence#186.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAS30638 standard; DNA; 32174 BP.
DNA encoding novel lung cancer antigen, Seq ID No WO200155300-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    MCACCIOL.
02-MG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 4; I
lery Match
10.3%; Score 28; DB 4; I
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           02-40G-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 4; I
LET_ --1 cimilarity 50.0%; Pred. No. 8.7e+02;
        10.3%; Score 28; DB 6;
63.2%; Pred. No. 7.8e+02;
                                                                                                                                                            28; DB 6;
No. 7.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.3%; Score 28; DB 4; 50.0%; Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                            10.3%; Score 28; DB 10; 60.5%; Pred. No. 8.3e+02;
                                                                                                                                                              Score
Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAS28700 standard; DNA; 32174 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DNA; 32174 BP
                                                                                                                                                                                                                                                                                                                                                                      06-NOV-2003.
(GENA-) GENAISSANCE PHARM INC.
                                                                                                      02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 1046
Query Match
Best Local Similarity
RESULT 1041
                                                          ABL70481 standard;
Chemically treated
WO200202807-A2.
                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 1043
                                                                                                                                                          Query Match
Best Local Similarity
RESULT 1042
                                                                                                                                                                                                                                                                                                                                                                                                                              Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 1050
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAL07491 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Si
RESULT 1047
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                                                                                                                                                                                                                                                         Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADC01147 standard; DNA; 16943 BP.
Strerohaemorragic B. coli 0157:H7-specific nucleic acid SEQ ID NO: 1191.
10-DEC-2002.
                                                                                                                                                    AAS46594 standard; DNA; 13469 BP.
Tumour suppressor gene derived chemically modified sequence #316.
WO200168912-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL08688 standard; cDNA; 21509 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 20546.
WO200171042-A2.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.3%; Score 28; DB 10; Length 17897; 66.7%; Pred. No. 7.1e+02;
                                                                                                                                                                                                                                                                                                     AAV74516 standard; DNA; 16397 BP.
Staphylococcus aureus contig SEQ ID #205.
EPP86519-A2.
30-JUL-1997.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
st Local Similarity 50.8%; Pred. No. 6.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 16943
                                                                                                10.3%; Score 28; DB 6; Length 10528; 48.2%; Pred. No. 5.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.3%; Score 28; DB 9; Length 16950; 49.3%; Pred. No. 6.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 17897
                                                                                                                                                                                                                                                       Length 13469;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.3%; Score 28; DB 4; Length 21509; 53.7%; Pred. No. 7.6e+02;
                    Human immune system associated gene SEQ ID NO: 308.
WO200200928-A2.
                                                                                                                                                                                                   Moderator.
(EPIG-2001.
(EPIG-) EPIGENOMICS AG.
(ery Match 10.3%; Score 28; DB 4; I 5ery Match aimilarity 56.5%; Pred. No. 6.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human metastasis associated gene SEQ ID NO: 175.
W0200177376-A2.
18-OCT-2001.
(EPIG-) BPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 28; DB 10;
Pred. No. 7.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.3%; Score 28; DB 10; 49.3%; Pred. No. 6.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACD19153 standard; DNA; 16950 BP.
E. coli 0157 unique DNA sequence OZID_166.
US2003023075-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Pretreated genomic DNA; 17897 BP. WO2003072821-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                RESULT 1037

ID ADB54177 standard, DNA, 17897 BP.
DE Pretreated genomic DNA region 101
PN WC0203072821-A2.
PD 04-SEP-2003.
PA (EPIG-) EPIGENOMICS AG.
  ABL32335 standard; DNA; 10528 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABL34622 standard; DNA; 23683 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.3%;
                                                      03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (BLAT) BLATTNER F R. (BURL) BURLAND V D. (PERN) PENNA N T. (PLUN) PLUNKETT G. (WELC) WELCH R.
                                                                                                                                                                                                                                                                                                                                       ...A-) HUMAN GENOM.
..rry Match
Best Local Similarity PRESULT 1035
ID ADC01147 stand*
DE Enterchaem*
PN JP2002**
PD 10**
                                                                                                                                                                                 ...ss
...2-A2.
...ac.) EPIGENOMICS
..ery Match
Best Local Similarity ...
RESULT 1034
ID AAV74516 stand?
DE Staphylocor
PN SEP78651°
PD 30-
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (UYTS-) UNIV TSUKUBA.
                                                                                                Query Match
Best Local Similarity
RESULT 1033
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity
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Best Local Similarity
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1D ACD1915:

DE B. CO1915:

PD 30-JAN.

PA (BURL/)

PA (PERN/)

PA (PERN/)

PA (PERN/)

PA (PERN/)

PA (PERN/)

PA (PERN/)
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PP PR

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AAK68883 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:23695.
WO200157182-A2.
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Human immune/haematopoietic antigen genomic sequence SEQ ID NO:30682.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAK72925 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:27737.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                      ABX59268 standard; cDNA; 32174 BP. cDNA encoding novel human musculoskeletal system antigen #1612. US2002147140-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SEQ ID
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                                                                                                      Length 32174;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADG41896 standard; DNA, 32174 BP.

Human respiratory system associated genomic DNA seq id 1134.
US2003215893-A1.
20-NOV-2003.
(HUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 10; Length 32174
st Local Similarity 50.0%; Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                      Length 32174;
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13-MAR-2003.

(HUMA.) HUMAN GENOME SCI INC.

(HAMA.) HUMAN GENOME SCI INC.

(Ery Match

"Anilarity 50.0%; Pred. No. 8.7e+02;
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(HUWA-) HUMAN GENOME SCI INC.
(HUWA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 4; I
Lery Match 10.3%; Pred. No. 9.3e+02;
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(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 4; Sery Match Augustanity 50.0%; Pred. No. 9.38+02;
                                                                                                                                          ESULT 1060.

ID ACA03401 standard; DNA; 32174 BP.

DE DNA encoding human lung cancer antigen HIPAA05.

PN US2002174454-A1.

PD 21-NOV-2002.

PA (ROSE/) ROSEN C A.

PA (RUBE/) RUBEN S M.

PA (BARA/) BARASH S C. 10 1%: Score 28; DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 28; DB 8;
Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB 12;
8.7e+02;
                                                                                                      Score 28; DB 6;
Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                      Score 28; DB 8;
Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADB96749 standard; DNA; 32174 BP.
Novel lung cancer antigen genomic DNA #20.
US2003049703-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          15-5AN-2004.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 10.3%; Score 28;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA; 32174 BP
                                                                                                        10.3%;
50.0%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                     10-OCT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
RESULT 1064
                                        (ROSE/) ROSEN C A.
(RUBI/) RUBIN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
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Best Local Similarity
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                                                                                                                            Local Similarity
                                                                                                                                                                                                                                                                                                                                            Local Similarity
US2002045230-A1.
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  Genomic sequence #540 encoding for novel human respiratory antigen
                                                                                                                                                                                                           D2-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
10.3%; Score 28; DB 4; Length 32174;
                                                                                                                                                                                                                                                                                                                                                                                    OZ-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 10.3%; Score 28; DB 4; Length 32174;
ery match 50.0%; Pred. No. 8.7e+02;
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(HUMA-) HUMAN GENOME SCI INC.
(HTMA-) HUMAN GENOME SCI INC.
ery Match
(ery Match 10.3%; Score 28; DB 5; Length 32174;
ery Match 20.0%; Pred. No. 8.7e+02;
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Human nervous system related polynucleotide SEQ ID NO 12690.
W0200159063-A2.
16-AUG-2001.
(HUMA) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 5; Length 32174; St Local Similarity 50.0%; Pred. No. 8.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 32174;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABA21505 standard; DNA; 32174 BP.
Human nervous system related polynucleotide SEQ ID NO 13836.
WO200159063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
10.3%; Score 28; DB 5; Length 32174;
Best Local Similarity 50.0%; Pred. No. 8.7e+02;
RESULT 1056
1D ABA19477 standard; DNA; 32174 BP.
DB Human nervous system related polynucleotide SEQ ID NO 11808
PN WO200159063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 32174
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                                                                                    Length 32174
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Human nervous system related polynucleotide SEQ ID NO 7996.
WOZOOLS9063-A2.
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Query Match
Best Local Similarity 50.0%; ...
RESULT 1053
ID ABR72131 standard; DNA; 32174 BP.
DE Human ovarian antigen #47 genomic sequence #1.
PD W0200155329-A2.
PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
PA (HUMA-) Total Color Sci DB 5; Leng
"...ch "...tv 50.0%; Pred. No. 8.7e+02;
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Human DNA for a novel foetal antigen, SEQ ID No 1818
WO200155312-A2.
                                                                                                                                                   AAS32655 standard; DNA; 32174 BP.
Human genomic DNA for novel endocrine antigen, SEQ
WO200155319-A2.
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16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
ery Match
10.3%; Score 28; DB 5; I
ery Match
10.3%; Pred. No. 8.7e+02;
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8.7e+02;
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                                                                                    10.3%; Score 28; DB 4; 50.0%; Pred. No. 8.7e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.3%; Score 28; 50.0%; Pred. No.
                                                                                                                                                                                                                                                                                                                        AAI62904 standard; DNA; 32174 BP.
Human genomic DNA SEQ ID NO 232.
WO200155449-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
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                                                                (HUMA-) HUMAN GENOME SCI INC.
                                                                                Query Match
Best Local Similarity
RESULT 1051
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Best Local Similarity
RESULT 1058
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Best Local Similarity
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Best Local Similarity
RESULT 1052
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 1055
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 1057
                        WO200155448-A1.
02-AUG-2001.
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Best Local Similarity 50.0%; Pred. No. 9.3e+02; RESULT 1077
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                                                                                             21-NOV-2002.
(HUMA-) HUMAN GENOME SCI INC.
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(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           30-JAN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       13-DEC-2001.
(AVAL-) AVALON PHARM.
                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
RESULT 1080
                                                                                                                                                                                                                                                                          Best Local Similarity RESULT 1079
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 1081
                                                                                                                                   Query Match
Best Local Similarity
RESULT 1078
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Best Local Similarity
RESULT 1084
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                                                                                                                                                                                                                                                                Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                NA DE
                                                          AAK71499 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:26311.
WO200157182-A2.
                                                                                                                                                                                          AAK63538 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:38350.
WO200157182-A2.
09-AUG-2001.
                                                                                                                                                                                                                                                                                                                        AAK69604 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:24416.
09-AUG-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAK80325 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:35137.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAK66361 standard; DNA; 38771 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:21173.
WC200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Secreted protein gene 264 genomic fragment HPMCJ84, SEQ ID NO:1517.
WO200277013-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABZ74036 standard; DNA; 38771 BP.
Secreted protein gene 165 genomic fragment HKACI79, SEQ ID NO:1183.
WC200277013-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABZ74376 standard; DNA; 38771 BP.
Secreted brotein gene 265 genomic fragment HPMCV30, SEQ ID NO:1523.
WO200277013-A2.
03-0CT-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              W.Z. 2003.
(BUMA-) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 8; Length 38771;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 10.3%; Score 28; DB 4; Length 38771;
ery match 50.0%; Pred. No. 9.3e+02;
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                                                                                                                                               Length 38771;
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                                                                                                        09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ELY Match 10.3%; Score 28; DB 4; I
(ELY Match 50.0%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (HUMA-) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 50.0%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                            10.3%; Score 28; DB 4; 1 50.0%; Pred. No. 9.3e+02;
               Score 28; DB 4;
Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                         (HUMA-) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 50.0%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.3%; Score 28; DB 4; 50.0%; Pred. No. 9.3e+02;
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03-OCT-2002.
(HUMA) HUMAN GENOME SCI INC.
(HUMA) HUMAN GENOME SCI INC.
10.3%; Score 28; DB 8; "ery Match 10.3%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADA44358 standard; DNA; 38771 BP.
Human secreted protein DNA SEQ ID 551.
WO2003000865-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABZ74370 standard; DNA; 38771 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
               10.3%;
50.0%;
                                                                                                                                                                                                                                                           (HUMA-) HUMAN GENOME SCI INC.
                                                                                     Best Local Similarity
RESULT 1075
DE ABZ74376 standard; DN
DE Secreted protein gene
PN W0200277013-A2.
PD 0-OCT-2002.
PA (HUMA-) HUMAN GENOME 8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 1074
ID ABZ74036 standard; DN
DE Secreted protein gene
PN WC200277013-A2.
PD 03-OCT-2002.
PA (HUMA-) HUMAN GENOME
           Query Match
Best Local Similarity
REGULT 1068
ID AAK71499 standard, DN
DE Human immune/haematop
PN WO20157182-A2.
PD 09-A016-2001.
PA (HUMA-) HUMAN GENOME
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 1072
ID AAK6636:
DE Human in
PN WC20015:
PD 09-AUG-:
PA (HUMA-)
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15-ARK-ZUU4.
(MORK/) MORELA D W.
(ENGE/) ENGELHARD E K.
10.3%; Score 28; DB 12; Length 96593;
                                                                                                                  Length 38771;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 38771;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 28; DB 10; Length 96593;
Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                                                                     Length 38771;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 51259;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 89328;
                                                                                                                                                                              ABZ67623 standard; DNA; 38771 BP.
Human secreted protein encoding genomic DNA SEQ ID NO 1146.
WO200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                                          Human secreted protein encoding genomic DNA SEQ ID NO 1464 W0200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human secreted protein encoding genomic DNA SEQ ID NO 1470 WO200277186-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL61995 standard; DNA; 89328 BP. Colon adenocarcinoma related gene sequence SEQ ID NO:332 WO200194629-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Mouse Blm carcinoma associated gene, SEQ ID NO:1403
WO2003057146-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Murine carcinoma associated (CA) nucleic acid #76 US2004072154-A1.
                   Human secreted protein-related DNA sequence #271 WO200292787-A2.
                                                                                                                                                                                                                                             03-OCT-2002.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 10.3%; Score 28; DB 10;
ery Match 50.0%; Pred. No. 9.38+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.3%; Score 28; DB 10; 50.0%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 28; DB 6; ]
Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 28; DB 9; ]
Pred. No. 1.3e+03;
                                                                                                               10.3%; Score 28; DB 10; 50.0%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.3%; Score 28; DB 10; 50.0%; Pred. No. 9.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         OCAUL-1997,
(DARW-) DARWIN MOLECULAR CORP.
(DARW-) DARWIN MOLECULAR CORP.
10.3%; Score 28; DB 2;
(ery Match 19rity 58.3%; Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAX83007 standard; DNA; 51259 BP.
Partial mouse WRN genomic sequence #3.
W09724435-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADB72623 standard; DNA; 96593 BP.
Mouse Blm gene.
WO2003008583-A2.
ADC20853 standard; DNA; 38771 BP
                                                                                                                                                                                                                                                                                                                                                                    ABZ67941 standard; DNA; 38771 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABZ67947 standard; DNA; 38771 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADA02885 standard; DNA; 96593 BP.
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ADP45593 standard; DNA; 147300 BP.
Human Rho family guanine-nucleotide exchange factor KIAA0861 gDNA.
WC2004047623-A2.
10-JUM-2040.
(SEQU-) SEQUENOM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
Query Match
Best Local Similarity 45.5%; Pred. No. 1.5e+03;
RESULT 1091
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 147300;
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                                                                                                                                    Length 96594;
                                                                                                                                                                                                                                                  28; DB 6; Length 110000;
No. 1.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                          DB 2; Length 116624;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 28; DB 6; Length 222930;
Pred. No. 1.7e+03;
                                                                                                                                                                                      Score 28; DB 2; Length 110000;
Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 27.8; DB 6; Length 354;
Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 311
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABK84349 standard; cDNA; 222930 BP.
Human cDNA differentially expressed in granulocytic cells
WO200228999-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human soft tissue sarcoma-upregulated DNA - SEQ ID 2720. W02004048938-A2.
10-JUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
10.3%; Score 28; DB 12; Length 1 St Local Similarity 48.2%; Pred. No. 1.5e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABL87180 standard; cDNA; 311 BP.
Human ovarian cancer related cDNA clone SEQ ID NO:10158
WO200192581-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABL85170 standard; cDNA; 354 BP.
Human ovarian cancer related cDNA clone SEQ ID NO:8148
WO200192581-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.2%; Score 27.8; DB 6; 54.4%; Pred. No. 1.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                          10.3%; Score 28; DB 2; I 52.6%; Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.3%; Score 28; DB 12; 50.0%; Pred. No. 1.5e+03;
                                                                                                                                    Score 28; DB 10;
Pred. No. 1.3e+03;
    58.3%; Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                      Score
Pred.
                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADC86530 standard; DNA; 141912 BP.
Human GPCR gene SEQ ID NO:983.
EP1270724-A2.
02-JAN-2003.
                                                                                                                                                                                                                                                                                                            AAV52850 standard; DNA; 116624 BP.
Human eyal gene contig 4405-9480.
WO9832849-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADQ19901 standard; DNA; 158811 BP
                                    ADC85364 standard; DNA; 96594 BP. Human Pap coding sequence.
WOX003045230-A2.
05-JUN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.2%;
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11-APR-2002.
(GENE-) GENE LOGIC INC.
10.3%;
                                                                                                                                    10.3%;
58.3%;
                                                                                                                                                                                            10.3%;
50.0%;
                                                                                                                                                                                                                                                    10.3%;
49.3%;
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(INSP ) INST PASTEUR.
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(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  06-DEC-2001.
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
RESULT 1092
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 1093
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Best Local Similarity
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Best Local Similarity
RESULT 1086
                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 1088
                                                                                                                                                      Best Local Similarity RESULT 1087
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Best Local Similarity
RESULT 1089
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Best Local S
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AAII0180 standard; DNA; 467 BP.

S Probe #113 for gene expression analysis in human cervical cell sample.

NO200157278-A2.

O9-AUG-2001.

A (MOLE) MOLECULAR DYNAMICS INC.

10.2%; Score 27.8; DB 4; Length 467;

Best Local Similarity 50.4%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAI31430 standard; DNA; 467 BP.
Probe #116 used to measure gene expression in human placenta sample.
WO200157272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Probe #108 for gene expression analysis in human heart cell sample.
WO200157274-A2.
                                                                                                                                                                                                                                                                                                                                                                                                ABX36295 standard; cDNA; 411 BP.
Bovine EST associated with lactation/muscle/fat deposition #1460.
US2002137139-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABX42784 standard; cDNA; 412 BP.
Bovine EST associated with lactation/muscle/fat deposition #7949.
US2002137139-A1.
                                                                                                                                                                              Length 388;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 411;
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Human foetal liver single exon nucleic acid probe #118.
M020017277-A2.
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(99-AUCLE-) MOLECULAR DYNAMICS INC.

(MOLE-) MOLECULAR DYNAMICS INC.

10.2%; Score 27.8; DB 4;

ery Match

10.2%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
10.2%; Score 27.8; DB 4;
ery Match
10.2%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          PD 09-AUG-2001.

PA (MOLE-) MOLECULAR DYNAMICS INC.

Query Match

Best Local Similarity 50.4%; Pred. No. 2.2e+02;

RESULT 1102
                                                                                                                                                                                10.2%; Score 27.8; DB 9; 49.0%; Pred. No. 2e+02;
                                                                                                                                                                                                                                                                                                                                      Score 27.8; DB 4;
Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 27.8; DB 8;
Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.2%; Score 27.8; DB 8; 51.2%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                         Human polymucleotide SEQ ID NO 11456. WO200164835-A2.
ACH16903 standard; cDNA; 388 BP.
Hwan adult heart cDNA #1217.
US2003073623 Al.
17-APR-2003.
(DRWA/) DRMANAC R T.
(LABA/) LABAT I.
(STAC/) STACHE-CRAIN B.
(DCCK/) DICKSON M C.
(JONE/) JONES L W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DNA; 467 BP.
                                                                                                                                                                                                                                                                                                                                         10.2%;
59.5%;
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51.2%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                        26-SEP-2002.
(BYAT/) BYATT J C.
(MATH) MATHIALAGAN N
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             26-SEP-2002.
(BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Local Similarity
                                                                                                                                                                                                                       RESULT 1097
ID AAI91396 standard;
                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
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ID ABA51813 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABA21642 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Local Similarity
                                                                                                                                                                                                   Best Local Similarity
                                                                                                                                                                                                                                                                                                                        (HYSE-) HYSEQ INC.
                                                                                                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT 1099
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AAK26016 standard; DNA; 478 BP.
Human bone marrow expressed single exon probe SEQ ID NO:
WO200157276-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADL37148 standard; DNA; 549 BP.
Human ovarian cancer DNA marker #11038.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human ovarian cancer DNA, 580 BP.
WO2001-0979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Human ovarian cancer DNA marker #4741.
WO200170979-A2.
                                                                                                                                                                                                        09-AUG-2001,
(MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADI71999 standard; DNA; 549 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            , MILLENNIUM , MILLENNIUM BEST Local Similarity RESULT 1121 ID BE UD43532 -
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    16-OCT-2003.
(PENN/) PENN S G.
(RANK/) RANK D R.
(HANZ/) HANZEL D K.
                                                                                                                                                                                                                                                          Best Local Similarity RESULT 1115
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity
RESULT 1118
                                                                                                                                                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 1116
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 1119
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 1120
                                                             09-AUG-200
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        15-NOV-200
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      27-SEP-200
                                                                                                                                                                                                                                                Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
                                                                                                  Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Probe #578 for gene expression analysis in human cervical cell sample. WO200157278-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAI31899 standard; DNA; 478 BP.
Probe #585 used to measure gene expression in human placenta sample.
WO200157272-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABA22093 standard; DNA; 478 BP.
Probe #559 for gene expression analysis in human heart cell sample.
WC200157274-A2.
(MOLE-) MODECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAI00122 standard; DNA; 467 BP.
Probe #113 used to measure gene expression in human breast sample.
WO200157270-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity 50.4%; Pred. No. 2.2e+02;
RESULT 1108
ABSO1026 standard; DNA; 467 BP.
DE Human genome-derived single exon probe from lung SEQ ID No 117.
               AAK25556 standard; DNA; 467 BP.
Human bone marrow expressed single exon probe SEQ ID NO: 113.
WO200157276-A2.
                                                                                                                                                                                                                                                                Length 467;
                                                                                                                 Length 467;
                                                                                                                                                                                                                                                                                                                                                                                                           10.2%; Score 27.8; DB 4; Length 467; 50.4%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 467;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (MOLE-) MOLECULAR DYNAMICS INC.
ry Match
t Local Similarity 69.1%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.2%; Score 27.8; DB 4; Length 478; 69.1%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 478
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 478
                                                                                                                                                                                     Human brain expressed single exon probe SEQ ID NO: 111 WO200157275-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human foetal liver single exon nucleic acid probe #597 WO200157277-A2.
                                                                 09-AUG-2001.
(WOLE-) MOLECULAR DYNAMICS INC.
(ery Match 10.2%; Score 27.8; DB 4;
                                                                                                                                                                                                                09-AUG-2001.
(WOLE-) MOLECULAR DYNAMICS INC.
(ery Match 10.2%; Score 27.8; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           15-NOV-2010.

(MOUE-) MOLECULAR DYNAMICS INC.

(MOLE-) MOLECULAR DYNAMICS INC.

10.2%; Score 27.8; DB 6;

LETY MATCH 50.4%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       MOJECTOR DYNAMICS INC.
(MOJEC) MOLECULAR DYNAMICS INC.
10.2%; Score 27.8; DB 4;
lery Match
10.2%; Score 27.8; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.2%; Score 27.8; DB 4; 69.1%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                            ABSISI27 standard; DNA; 467 BP.
Human liver single exon probe, SEQ ID No 117.
WO200157273-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                    09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                 AAK00120 standard; DNA; 467 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABA52292 standard; DNA; 478 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT 1109

ID AA110645 standard; DNA; 478

DE Probe #578 for gene expressi

PN WOZDO157278-A2.

PD 09-AUG-2001.

PA (MOLE-) MOLECULAR DYNAMICS T
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 1113
                                                                                                           Query Match
Best Local Similarity
RESULT 1105
                                                                                                                                                                                                                                                                           Best Local Similarity
RESULT 1106
                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 1107
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
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Best Local Similarity
RESULT 1111
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    09-AUG-2003
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 RESULT 1104
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AA100575 standard; DNA; 478 BP.
Probe #566 used to measure gene expression in human breast sample.
WO200157270-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity 10:18; Pred. No. 2.2e+02; RESULT 1117
ID ABSOOF57 standard; DNA; 478 BP.
BB Human genome-derived single exon probe from lung SEQ ID No 588.
PN WO200186003-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 520;
                             Length 478;
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                                                                                                                                                                                                                                   Length 478;
                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 478;
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                                                                          RESULT 1114
ID AAKOOS67 standard; DNA; 478 BP.
DE Human brain expressed single exon probe SEQ ID NO:
PN WO200157275-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.2%; Score 27.8; DB 12; 54.4%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                                                                                                                                                     Weacter

09-ADG-2001.

(MOLE-) MOLECULAR DYNAMICS INC.

(MOLE-) MOLECULAR DYNAMICS INC.

10.2%; Score 27.8; DB 4;

ery Match

10.2%; Score 27.8; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (MOLE-) MOLECULAR DYNAMICS INC.
10.2%; Score 27.8; DB 6;
it Local Similarity 69.1%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.2%; Score 27.8; DB 5;
17-7-71 Similarity 54.4%; Pred. No. 2.3e+02;
(MOLE-) MOLECULAR DYNAMICS INC.

10.2%; Score 27.8; DB 4;

t Local Similarity 69.1%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
ery Match 10.2%; Score 27.8; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 10.2%; Score 27.8; DB 5;
er Tocal Similarity 54.4%; Pred. No. 2.3e+02;
                                                                                                                                                                                                                                 10.2%; Score 27.8; DB 4; 69.1%; Pred. No. 2.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.2%; Score 27.8; DB 5; 54.4%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                      ABS25603 standard; DNA; 478 BP.
Human liver single exon probe, SEQ ID No 593.
WO200157273-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACH73507 standard; DNA; 520 BP.
Human genome derived single exon probe #6702.
US2003194704-Al.
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Length 1276;

Length 1342

Length 1613;

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ADPO4618 standard; cDNA; 1707 BP.
Sea squirt cDNA with tissue specific expression in development Seq 213.
JP2004057129-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAQ25532 standard; DNA; 1648 BP.
Sequence of genomic clone contg. the entire Histidine-rich protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match 10.2%; Score 27.8; DB 10; Length 1803; Best Local Similarity 53.2%; Pred. No. 3.66+02; RESULT 1138
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAH28330 standard; DNA; 2370 BP.
Nucleotide sequence of a bacterial surface array protein (SAP)
RESULT 1131

ID ABRITYO33 standard; CDNA; 1276 BP.

E CDNA encoding human G-protein coupled receptor GCREC-57.

PN WO200279448-A2.
                                                                                                                                                        Best LUCL...
SULT 1132

D ADC86520 standard; DNA; L3-2.
E Human GPCR gene SEQ ID NO:773.
E Human GPCR gene SEQ ID NO:773.
PD 02-JAN-200-3.
PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
PA (NAAD-) CENT ADVANCED SCI & TECHNOLOGY.
PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY.

TO MATCH 10.2%; Score 27.8; DB 10; Ler

"Match 1-1-rity 47.9%; Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                            RESULT 1133

TD ABX71384 standard; cDNA; 1613 BP.

DB Human testes-derived cDNA from clone DKFZphtes3_297.

PN W0200112659-A2.

TO TO THE TOWN PROJECT.

TO THE TOWN PROJECT.
                                                                           MCZCZ-2002.
10-OCT-2002.
(INOY-) INCYTE GENOMICS INC.
(ELY MATCh 10.2%; Score 27.8; DB 10;
(ELY MATCh 47.9%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.2%; Score 27.8; DB 12; 51.2%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                       22-FEB-2001.
(GEHU-) GERMAN HUMAN GENOME PROJECT.
ery Match 10.2%; Score 27.8; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           PD 26-MAY-1992.
PA (SLOK ) SLOAN KETTERING INST CANCER.
Query Match 10.2%; Score 27.8; DB 2;
Best Local Similarity 54.4%; Pred. No. 3.5e+02;
RESULT 1135
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.2%; Score 27.8; DB 6; 65.1%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.2%; Score 27.8; DB 4; 53.2%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABS76421 standard; cDNA; 2611 BP. cDNA encoding human ovarian cancer marker M445. WO200271928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human cDNA encoding clone TESTI20049990.

BPI308459-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  26-FEB-2004.
(KAGA-) KAGAKU GIJUTSU SHINKO JIGYODAN.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               RESULT 1136
ID ABQ67836 standard; DNA; 1716 BP.
DE Listeria innocua DNA sequence #638.
PN WO200228891-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              07-MAY-2003.
(HELI-) HELIX RES INST.
(REAS-) RES ASSOC BIOTECHNOLOGY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  12-JUL-2001.
(BIOS-) BIOSITE DIAGNOSTICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 11-APR-2002.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 1137
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (HiskP) gene.
US5116965-A.
                                                                                                                                    Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                RESULT 1134
1D AAQ2553.
DE Sequence
DE (HisRP)
PN US1169
PD 26-MAY-
PA (SLOK )
                                                                                                                                                 Best Local Similarity --- RESULT 1123

ID ARQ45591 standard; DNA; 602 BP.

ID ARQ45591 standard; DNA; 602 BP.

DE Oligomucleotide for detecting cytosine methylation SEQ ID NO 32182.

PN W0200218632-A2.

PA (EPIG-)

Query Match

Toral Similarity 53.2%; Pred. No. 2.46+02;
RESULT 1122

ID ABGA5590 standard; DNA; 602 BP.

DE Oligonucleotide for detecting cytosine methylation SEQ ID NO 32181.

PN W0200218632-A2.

PD 07-MAR-2002.

PD 07-MAR-2002.

PA (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 27.8; DB 8; Length 1194; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAX20759 standard; DNA; 656 BP.
Polynuclectide sequence from the genome of Treponema pallidum WO9859034-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.2%; Score 27.8; DB 6; Length 1029; 57.5%; Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 1099;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                30.DEC-1998.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
10.2%; Score 27.8; DB 2; Length 656;
ery Match
34.4%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 701;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 682
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 609
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAZ77517 standard; CDNA; 701 BP.
Human ovarian tumor cDNA library derived BST fragment
DE19817557-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                cDNA; 682 BP.
human diagnostic protein #5377
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.2%; Score 27.8; DB 3; 50.4%; Pred. No. 3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.2%; Score 27.8; DB 5; 53.2%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                     22-FEB-2001.
(BETH-) BETH ISRAEL DEACONESS MEDICAL CENT.
ery Match
10.2%; Score 27.8; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.2%; Score 27.8; DB 2; 48.4%; Pred. No. 2.5e+02;
                                                                                                             AG.
10.2%; Score 27.8; DB 6;
53.2%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                          AAF76057 standard; DNA; 609 BP.
Enterococcus faecium vanXD gene, SEQ ID NO:39.
WO200112803-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity 48.4%; Pred. No. 2.5e4-RESULT 1128
ID ABNG726 standard; DNA; 1029 BP.
DE Streptococcus polynucleotide SEQ ID NO 2365.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAC67659 standard; cDNA; 1099 BP.
Human secreted protein cDNA sequence #29-
WO200058355-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    GENOMFORSCHUNG MBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Prokaryotic essential gene #9654 WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DNA; 1194 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              WOZUGE...
03-OCT-2002.
(ELIT-) ELITRA PHARM INC.
10.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   02-MAY-2002.
(CHIR-) CHIRON SPA.
(GENO-) INST GENOMIC RES.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        21-OCT-1999.
(META-) METAGEN GES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1125
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAS69573 standard;
DNA encoding novel
WO200175067-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
RESULT 1129
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1130
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
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RESULT 1127
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ACA27997 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11-OCT-2001
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Query Match

Query Match

Query Match

Length 1707

Length 1648

Length 1716;

Query Match

Query Match

Length 2370;

PN

PP PP

ID DE PN PD

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AAS45367 standard; DNA; 8842 BP.
Chemically pretreated complementary DNA associated with cell cycle #36.
20-SEP-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABL70338 standard; DNA; 6226 BP.
Chemically treated cell signalling DNA sequence complementary to#114.
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Signal transduction associated gene modified complementary DNA \#117.80200200926-A2.
                                                                                                                                                                                                                                                          ABZS8670 standard; DNA; S829 BP.
Human KCNMA gene related DNA (GenBank Identifier No. GI#7914977).
WO200299058-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAS46618 standard; DNA; 6863 BP.

Humour suppressor gene derived chemically modified sequence #340.
20-SEP-2001.
EPIGENCMICS AG.
10.2%; Score 27.8; DB 4; Length 6863;
ELocal Similarity 50.4%; Pred. No. 5.8e+02;
                                                                                                                         ID NO:4
                                           Score 27.8; DB 10; Length 4933; Pred. No. 5.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 6115;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 5945;
                                                                                                                                                                                                                                                                                                                                                             Length 5829;
                                                                                                                                                                                                       Length 5781
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 6226,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity 62.0%; Pred. No. 5.6e+02; RESULT 1155
ID AAS61293 standard; DNA; 6226 BP.
DE Human gene regulation-associated gene oligonucleotide #248.
PN WO200177375-A2.
                                                                                                  AAF76022 standard; DNA; 5781 BP.
E. faecium VanD vancomycin resistance gene cluster, SEQ
WO200112803-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                           730
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADF90711 standard; DNA; 6115 BP.
Human hepatic-fibrosis disease marker SEQ ID 173
JP2003259877-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.2%; Score 27.8; DB 10; 51.2%; Pred. No. 5.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.2%; Score 27.8; DB 6; 62.0%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                             Score 27.8; DB 9;
Pred. No. 5.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 27.8; DB 6;
Pred. No. 5.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.2%; Score 27.8; DB 6; 62.0%; Pred. No. 5.6e+02;
                                                                                                                                                                                                   10.2%; Score 27.8; DB 4; 52.1%; Pred. No. 5.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 27.8; DB 6;
Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                ABL32757 standard; DNA; 5945 BP.
Human immune system associated gene SEQ ID NO:
03-0784-2002.
                                                                                                                                                            (BETH-) BETH ISRAEL DEACONESS MEDICAL CENT.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABK31391 standard; DNA; 6226 BP.
                                           10.2%;
47.0%;
                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity 57.5%;
RESULT 1151
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WO2002.
10-J0H-2002.
(EPIG-) EPIGENOMICS AG.
10.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          OF-DATE AG. 10.2%; try Match 10.2%; tr Local Similarity 51.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      16-SEP-2003.
(SUMU ) SUMITOMO SEIYAKU KK.
    (GEHO ) GEN HOSPITAL CORP. (FARB ) BAYER AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 18-OCT-2001.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                    12-DEC-2002.
(EXEL-) EXELIXIS INC.
                                                         Best Local Similarity
RESULT 1149
                                                                                                                                                                                                               Best Local Similarity RESULT 1150
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 1154
ID ABL70338 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 1156
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
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                                           Query Match
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                                                                                                                                                                                                                                                                                                                                                                                            AEL49392 standard; DNA; 2872 BP. Whuman polynucleotide associated with DNA replication SEQ ID NO 92. W0200177377-A2. 18-OCT-2001.
                                                                                                                                                                                                                                                      Chemically treated DNA repair gene fragment complementary to#69 WO200181622-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAFBE149 standard; DNA; 4790 BP.
DNA encoding a dishevelled associated kinase (DAK) isoform DAKb
WO200125408-A1.
                                                                          ABL29192 standard; DNA, 2660 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 39049.
WQ20011042-A2.
Z7-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         27-NOV-2002.
(BIOW-) BIOWINDOW GENE DEV INC SHANGHAI.
ery Match
-- Tocal Similarity 53.2%; Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 4933;
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                      Length 2611;
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Pred. No. 4.7e+02;
                                                                                                                                                                              10.2%; Score 27.8; DB 4; Length 2660; 59.5%; Pred. No. 4.1e+02;
                                                                                                                                                                                                                                                                                                                                        Length 2872;
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50.4%; Pred. No. 5.1e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity 59.5%; Pred. No. 4.2e+02; RESULT 1143
ID ADD01140 standard; CDNA, 3190 BP.
DE Human signal transducer kinase 22.55 encoding cDNA #SEQ PN CN1381571-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL32171 standard; DNA; 3973 BP.
Human immune system associated gene SEQ ID NO: 144
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.2%; Score 27.8; DB 10; 47.0%; Pred. No. 5.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 27.8; DB 12;
Pred. No. 4.9e+02;
(MILL-) MILLENNIUM PHARM INC.

17.8; Score 27.8; DB 6; t Local Similarity 50.4%; Pred. No. 4.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 27.8; DB 6;
Pred. No. 4.2e+02;
                                                                                                                                                                                                                                                                                                                                      Score 27.8; DB 6;
Pred. No. 4.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADJ67448 standard; DNA; 4257 BP.
Human ovarian specific gene SEQ ID NO:162.
WO2004013311-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADD48702 standard; DNA; 4933 BP.
Rat gene M64793, SEQ ID NO 14411.
W02003016475-A2.
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ADD47239 standard; DNA; 4933 BP.
Rat gene M64793, SEQ ID NO 12933.
WC2003016475-A2.
27-FEB-2003.
                                                                                                                                                                                                                                      DNA; 2872 BP.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                1 (EPIC-1 EPIGENOMICS AG.
Query Match 10.2%;
Pact Local Similarity 59.5%;
                                                                                                                                                                                                                                                                                                              (EPIG-) EPIGENOMICS AG,
ry Match
t Local Similarity 59.5%;
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(GEHO ) GEN HOSPITAL CORP.
(FARB ) BAYER AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      03-JAN-2002.
(EPIG-) BPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (REGC ) UNIV CALIFORNIA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Ja.
...GC ) UNIV CALIFO.
...GC ) UNIV CALIFO.
...Ty Match
Best Local Similarity 5.
RESULT 1147
ID ADD48702 standar
DE Rat gene M647
PN WO200301f
PN GT FEP
PA (CT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  12-FEB-2004.
(DIAD-) DIADEXUS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
RESULT 1142
                  Query Match
Best Local Similarity
RESULT 1140
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 1144
                                                                                                                                                                                             Best Local Similarity RESULT 1141
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                                                                                                                                                                                                                                      ABL92329 standard;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
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Length 13894;

Length 14041;

Length 14861;

Length 14861;

Length 14861

Length 15732

Length 15732

Length 16920;

4 ;

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Chemically pretreated complementary DNA associated with cell cycle #47. WO200168911-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAK69093 standard; DNA; 16920 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:23905.
WO200157182-A2.
         27-MAR-2003.
( BCGEA-) EGEA BIOSCIENCES INC.
Query Match
Best Local Similarity 49.7%; Pred. No. 7.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABK28234 standard; DNA; 15732 BP.
DNA transcription associated complementary genomic DNA WO200192565-A2.
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Human gene regulation-associated gene oligonucleotide
WO200177375-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                           Human immune system associated gene SEQ ID NO: 1167
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DNA; 14861 BP.
cell signalling DNA sequence#222
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Enterococcus faecalis genome contig SEQ ID NO:128
WO9850555-A2.
12-NOV-1998.
                                                                                                                                                                                              Score 27.8; DB 12;
Pred. No. 7.5e+02;
                                                                                                                                                                                                                                                                                                                                                       Score 27.8; DB 4;
Pred. No. 7.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 27.8; DB 6;
Pred. No. 7.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 27.8; DB 6;
Pred. No. 7.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 27.8; DB 6;
Pred. No. 7.6e+02;
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Pred. No. 7.6e+02;
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Pred. No. 7.8e+02;
                                                                                     Mycoplasma gentalium functional gene fragment US20040397-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.2%; Score 27.8; DB 57.5%; Pred. No. 8e+02;
                                                                                                                                                                                                                                                          DNA; 14041 BP.
                                                                                                                                                                                                                                                                                                                                                                                                               DNA; 14861 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DNA; 15732 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAX13065 standard; DNA; 32768 BP
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10-JAN-2002.

(EPIG-) EPIGENOMICS AG.

10.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    WOZUGECCO.
20-SED-2001.
(EPIG-) EPIGENOMICS AG.
10.2%;
... Match ... arity 57.5%;
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06-DEC-2001.
(EPIG-) EPIGENOMICS AG.
10.2%;
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(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                     10.2%;
18.3%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                         WOZOGETO
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.2%;
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WO200146463-A2.
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                                                                                                                                                                                                                  Best Local Similarity
RESULT 1168
ID AAH48024 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL70553 standard;
Chemically treated
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1173
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 1171
ID AAS61202 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 1172
                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                 ABL33194 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAS45389 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
                                                                               RESULT 1167
ID ADN48952 standard;
                                                                                                                                                            01-APR-2004.
(EVAN/) EVANS G A.
                                                                                                                                                                                                                                                                                                              28-JUN-2001.
(BAXT ) BAXTER AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                    AAS45377 standard; DNA; 9091 BP.
Chemically pretreated complementary DNA associated with cell cycle #41.
WO200168911-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAK69092 standard; DNA; 9454 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:23904.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACC69147 standard; DNA; 13894 BP.
M. genitalium polypeptide chaperone gene cassette DNA SEQ ID NO:15.
WO2003025145-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 13894;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 11029
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        77 1163
ABZ68481 standard; DNA; 11029 BP.
Nucleotide sequence of the genome of West Nile virus IS-98-ST1
WO200281511-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 10254;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 9454;
                             Length 8842;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 9091;
                                                                                                                                                                                                                                                                                                                                   Length 9091;
                                                                                                                                                                                Length 8842
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABV74821 standard; DNA; 11029 BP.
West Nile virus strain NY99-flamingo 382-99 complete genome.
WO200281741-A2.
                                                                                                       #39.
                                                                                                                                                                                                                                                                                                                                                                                           ABK28214 standard; DNA; 9091 BP.
DNA transcription associated complementary genomic DNA #44
WO200192565-A2.
                                                                           ABK20204 standard; DNA; 8842 BP.
DNA transcription associated complementary genomic DNA
WO200192565-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABQ67046 standard; DNA; 10254 BP.
Human angiogenesis associated polynucleotide SEQ ID NO
WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.2%; Score 27.8; DB 10; 57.5%; Pred. No. 6.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.2%; Score 27.8; DB 10; 49.7%; Pred. No. 7.5e+02;
                                                                                                                                               06-DEC-2001.
(EPIG-) EPIGENOMICS AG.
2ry Match 10.2%; Score 27.8; DB 6;
rran Similarity 55.8%; Pred. No. 6.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AG.
10.2%; Score 27.8; DB 6;
57.5%; Pred. No. 6.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AG.
10.2%; Score 27.8; DB 6;
53.2%; Pred. No. 6.7e+02;
                                                                                                                                                                                                                                                                                                                                     Score 27.8; DB 4;
Pred. No. 6.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (HUMAN) HUMAN GENOME SCI INC.
10.2%; Score 27.8; DB 4;
t Local Similarity 57.5%; Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.2%; Score 27.8; DB 8; 57.5%; Pred. No. 6.9e+02;
                             Score 27.8; DB 4;
Pred. No. 6.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Mycoplasma genitalium chaperone gene.
US2003138777-A1.
24-JUL-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAD63517 standard; DNA; 13894 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                17-OCT-2002.
(INSP ) INST PASTEUR.
(KIMR-) KIMRON VETERINARY INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       17-OCT-2002.
(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
(EPIG-) EPIGENOMICS AG.
10.2%;
rronal Similarity 55.8%;
                                                                                                                                                                                                                                                                                                    20-SEP-2001.
(EPIG.) EPIGENOMICS AG.
197 Match 10.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               13-JUN-2002.
(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 1165
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 1166
                           Query Match
Best Local Similarity
RESULT 1158
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 1161
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Best Local Similarity
RESULT 1164
                                                                                                                                                                                                Best Local Similarity RESULT 1159
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RESULT 1160
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (EVAN/) EVANS G A.
                                                                                                                                             06-DEC-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                       06-DEC-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          09-AUG-200
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DE DE PN PD PD PA
                                                                                                                                                                                                                                                                                                                                            10.2%; Score 27.8; DB 10; Length 50000; 62.0%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.2%; Score 27.8; DB 10; Length 54355; 57.5%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.2%; Score 27.8; DB 10; Length 54355; 57.5%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 10; Length 66804;
                                                                                                                                                                                                                                                                                                                                                                                                                                               17-20-20-3.

17-20-20-3.

(SAGR-) SAGRES DISCOVERY.

6ry Match

10.2%; Score 27.8; DB 9; Length 54355;

ery Match

57.5%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 54355;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABK87050 standard; cDNA; 66804 BP.

Human transporter protein genomic DNA.

uery Match

10.2%; Score 27.8; DB 6; Length 66804;

est Local Similarity 54.4%; Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.2%; Score 27.8; DB 6; Length 81905; 65.1%; Pred. No. 1.4e+03;
                                                                                                                                                                                                  Length 32768;
                 10.2%; Score 27.8; DB 2; Length 32768; 49.7%; Pred. No. 1e+03;
                                                                                                                                                                                                                                                                      Human mammalian target of rapamycin genomic fragment WO2003048360-A1.
12-JUN-2003.
(NEWI-) NEW IND RES ORG.
                                                                                                                                                                                                                                                                                                                                                                                                                  Mouse Morf carcinoma associated gene, SEQ ID NO:1271 WO2003057146-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADM74348 standard; DNA; 54355 BP.
Murine carcinoma associated (CA) nucleic acid #10.
US2004072154-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.2%; Score 27.8; DB 12; 57.5%; Pred. No. 1.2e+03;
                                                                                                                                                                                                  10.2%; Score 27.8; DB 6; 49.7%; Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match Form 10.2%; Score 27.8; DB 6; Best Local Similarity 54.4%; Pred. No. 1.3e+03; ID ADG88330 Standard; DNA, 66804 BP.

DB Human transporter protein genomic DNA.

Query Match 10.2%; Score 27.8; DB 10 Best Local Similarity 54.4%; Pred. No. 1.3e+03; RESULT 1184
                                                                  ABS98860 standard; DNA; 32768 BP.
Enterococcus faecalis contig sequence #128.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABQ69244 standard; DNA; 81905 BP.
Listeria innocua DNA sequence #683.
WO200228891-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADB72491 standard; DNA; 54355 BP.
Mouse Morf gene.
WC2003008583-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity 57.5%; Pred
RESULT 1180
ID ADC85233 standard; DNA; 54355 BP.
                                                                                                                                                                                                                                                    ADC87687 standard, DNA, 50000 BP.
                                                                                                                                                                                                                                                                                                                                                                                                 ADA02753 standard; DNA; 54355 BP.
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(INSP ) INST PASTEUR.
(CNRS ) CNRS CENT NAT RECH SCI.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Mouse Morf genomic sequence.
WG22033045230-A2.
OG-JUN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    30-JAN-2003.
(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        15-APR-2004.
(MORR/) MORRIS D W.
(ENGE/) ENGELHARD E K.
                                                                                                                                                                                                                                                                                                                                                                                                             Carc.

J57146-A2.

ULD-2003.

(SAGR-) SAGRES DISC.

Query Match
Best Local Similarity PRESULT 1179
ID ADB72491 standar
DE Mouse Morf
PN WC20030^C
PA
                                                                                                      US2002120116-A1.
29-AUG-2002.
(KUNS) KUNSCH C A.
(DILLL) DILLON P J.
(BARA/) BARASH S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1182
                            Best Local Similarity RESULT 1176
                                                                                                                                                                                                                 Best Local Similarity
                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 1178
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Best Local Similarity
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                  Query Match
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ID ADM74340
DE Murine OPN US20040
PD 15-APR-',
PA (MORR/)
PA (ENGE/)
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Alloiococcus otitis antigenic protein encoding DNA SEQ ID NO:4237. WO2003048304-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 03-JUL-2003.
(INCY-) INCYTE GENOMICS INC.
(ENY Match 10.2%; Score 27.8; DB 10; Length 214019;
(F. 7.2.2) Similarity 69.1%; Pred. No. 1.96+03;
                                                                                                                                                                                                                                                                                                                                           Length 167163;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 10; Length 170170;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.2%; Score 27.8; DB 11; Length 276820; 50.4%; Pred. No. 2.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.2%; Score 27.8; DB 12; Length 262090; 51.2%; Pred. No. 2e+03;
                                                                                                                                                              Length 82689;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 282;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 294;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADL13809 standard; DNA; 214019 BP.
Osteoarthritis-associated polymorphic nucleotide #341.
WO2003054166-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                   Osteoarthritis-associated polymorphic nucleotide #175
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADQS9207 standard; DNA; 262090 BP.
MSI-H carcinoma genomic DNA sequence SEQ ID NO:44.
KR2004008012-A.
                                                                                                                                                                                                                                                                                     O2-OCT-2003.
A (SAGR-) SAGRES DISCOVERY.
Query Match 10.2%; Score 27.8; DB 10;
Query Match 52.1%; Pred. No. 1.8e+03;
                                                                                                                                                              10.2%; Score 27.8; DB 6; 65.1%; Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.1%; Score 27.6; DB 5; 50.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          23.AUG-2001.
1 (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Query Match
10.1%; Score 27.6; DB 5;
uest Local Similarity 50.0%; Pred. No. 2.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (INCY-) INCYTE GENOMICS INC.
Query Match
10.24; Score 27.8; DB 10
Best Local Similarity 49.74; Pred. No. 1.8e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABN59670 standard; cDNA; 282 BP.

E Human prostate expression marker cDNA 59661.

N W0200160860-A2.

23-W02001.

A (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity 50.0%; Pred. No. 2.1e+
RESULT 1192

ID ABV59998 standard; CDNA; 294 BP.

BE Human prostate expression marker CDNA 59989.

PN WG200160860-A2.

PD 33-AUG-2001.

PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC
RESULT 1185

ID ABQ67198 standard; DNA; 82689 BP.
DE Listeria innocua plasmid DNA sequence.
PN W020022891-A2.
PD 11-APR-2002.
PA (INSP ) INST PASTEUR.
PA (CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                                      Best Local Similarity 52.1%; Pred. RESULT 1187
ID ADL13643 standard; DNA, 170170 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DNA; 276820 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       17-APR-2003.
(GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         T 1193
ADB10825 standard; DNA; 417 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 1190
ID ADPTS188 standard; DI
DE Human ADAWTS2 gene.
PN WO2003031594-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 1188
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
RESULT 1189
                                                                                                                                                                            Best Local Similarity RESULT 1186
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       28-JAN-2004.
(KIMH/) KIM H G.
(KIMN/) KIM N G.
(LEEJ/) LEE J S.
(RHEE/) RHEE H S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                    WO2003054166-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               12-JUN-2003
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                                                                                                                                                                 Query Match
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Query Match

Query Match

Query Match

RESULT 1199

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AAI15582 standard; DNA; 448 BP.
Probe #5515 for gene expression analysis in human cervical cell sample.
WO200157278-A2.
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Probe #5775 used to measure gene expression in human placenta sample.
WO200157272-A2.
                                                                                                                     SEQ ID No 3167
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 1208
ID AAK31184 standard; DNA; 448 BP.
DE Human bone marrow expressed single exon probe SEQ ID NO: 5741
PN W0200157276-A2.
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Length 436;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 448,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABA27005 standard; DNA; 448 BP.
Probe #5471 for gene expression analysis in human heart
WO200157274-A2.
                                                                                                                                                                                                                                             Length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABAST534 standard; DNA; 448 BP.
Human foetal liver single exon nucleic acid probe #5839
WO200157277-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAKO5581 standard; DNA; 448 BP.

Human brain expressed single exon probe SEQ ID NO:
WO200157275-A2.
09-AUG-2001.
WOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABS05936 standard, DNA, 448 BP.
Human genome-derived single exon probe from lung
WO200186003-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         IN-CAULTINOV-2001.

(MOLE-), MOLECULAR DYNAMICS INC.

10.1%; Score 27.6; DB 6;
                                                                                     ABS03176 standard; DNA; 436 BP.
Human genome-derived single exon probe from lung
WO200186003-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MCACCAOOL.
09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
10.1%; Score 27.6; DB 4;
lery Match 10.1%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                     09-AUG-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
ery Match 10.1%; Score 27.6; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WOACCE...

09-AUG-2001.

(MOLE-) MOLECULAR DYNAMICS INC.

(MOLE-) MOLECULAR DYNAMICS INC.

10.1%; Score 27.6; DB 4;

10.7 Match

10.7 Matc
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (MOLE-) MOLECULAR DYNAMICS INC.

ry Match 10.1%; Score 27.6; DB 4;

t Local Similarity 55.1%; Pred. No. 2.5e+02;
                                                                                                                                                                       15.00-2001.
(MOLE-) MOLECULAR DYNAMICS INC.
(MOLE-) MOLECULAR DYNAMICS INC.
10.1%; Score 27.6; DB 6;
ery Match
10.1%; Soore 27.6; DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.1%; Score 27.6; DB 4; 55.1%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.1%; Score 27.6; DB 4; 55.1%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (MOLE-) MOLECULAR DYNAMICS INC.

10.1%; Score 27.6; DB 4;

t Local Similarity 55.1%; Pred. No. 2.5e+02;
Score 27.6; DB 5;
Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human liver single exon probe, SEQ ID No WO200157273-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DNA; 448 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (MOLE-) MOLECULAR DYNAMICS INC.
   10.1%;
53.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity
RESULT 1205
Query Match
Best Local Similarity
RESULT 1203
                                                                                                                                                                                                                                                             Best_Local Similarity
RESULT 1204
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABS30865 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            09-AUG-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      09-AUG-2001
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                                                                        AAI13242 standard; DNA; 436 BP.
Probe #3175 for gene expression analysis in human cervical cell sample.
WO200157278-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                T 1196
AA134594 standard; DNA; 436 BP.
Probe #3280 used to measure gene expression in human placenta sample.
09-AUG-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABA24706 standard; DNA; 436 BP.
Probe #3172 for gene expression analysis in human heart cell sample.
WO200157274-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAI03145 standard; DNA; 436 BP.
Probe #3136 used to measure gene expression in human breast sample.
WO200157270-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAK29665 standard; DNA; 436 BP.
Human bone marrow expressed single exon probe SEQ ID NO: 3222
WO200157276-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     436;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 436;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 436
                                  Length 417;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 436
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 436
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 436
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 436
                                                                                                                                                                                                                                                                                Length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length
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                                                                                                                                                                                                                                                                                                                                                               Assistant DNA; 436 BP.
Human foetal liver single exon nucleic acid probe #3244 WO200157277-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABA44498 standard; DNA; 436 BP.
Human breast cell single exon nucleic acid probe #3193
WO200157271-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        WLZVOLLO.

MOS-ACOLO.

(MOLE-) MOLECULAR DYNAMICS INC.

(MOLE-) MOLECULAR DYNAMICS INC.

Ery Match 10.1%; Score 27.6; DB 4;

ery Match 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                       09-AUG-2001.
(WOLE-) MOLECULAR DYNAMICS INC.
(WOLE-) MOLECULAR DYNAMICS INC.
(ery Match 10.1%; Score 27.6; DB 4;
(ery Match 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (MOLE-) MOLECULAR DYNAMICS INC.
ry Match 10.1%; Score 27.6; DB 4;
t Local Similarity 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              09-4001.
(WOLE-) MOLECULAR DYNAMICS INC.
(WOLE-) MOLECULAR DYNAMICS INC.
ery Match 10.1%; Score 27.6; DB 4;
ery Match 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10.1%; Score 27.6; DB 4; 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (MOLE-) MOLECULAR DYNAMICS INC.

ry Match

t Local Similarity 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (MOLE-) MOLECULAR DYNAMICS INC.
ry Match
t Local Similarity 53.8%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (MOLE-) MOLECULAR DYNAMICS INC.
ry Match
t Local Similarity 53.8%; Pred. No. 2.5e+02;
                                  10.1%; Score 27.6; DB 9; 50.0%; Pred. No. 2.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human liver single exon probe, SEQ ID No 3254 WO200157273-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          (MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (MOLE-) MOLECULAR DYNAMICS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DNA; 436 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABS28264 standard; DNA; 436 BP.
         (AMHP ) WYETH HOLDINGS CORP.
                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 1195
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 1197
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
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Best Local Similarity
RESULT 1201
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
                                     Query Match
Best Local Similarity
RESULT 1194
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAK03212 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            09-AUG-2001
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sample.

cell

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26-OCT-2000.
(UTAH ) UNIV UTAH RES FOUND.
(UYYA ) UNIV YALE.
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Best Local Si
RESULT 1227
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
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                                                                                                                                                 Query Match
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S. epidermidis open reading frame nucleotide sequence SEQ ID NO:2997.
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  RESULT 1213
L ARQSB582 standard; cDNA; 474 BP.
DE Human colon cancer related nucleotide sequence SEQ ID NO:2277
PN WO200229086-A2.
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(FROT-) PROTEIN DESIGN LABS INC.

(ery Match 10.1%; Score 27.6; DB 12; Length 481; Score 37.6; DB 12; Length 481; Score 37.6; DB 12; Length 481; Score 37.6; DB 12; Length 481; Length 4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.1%; Score 27.6; DB 12; Length 584; 50.0%; Pred. No. 2.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 472;
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PA (ZYMO) ZYMOGENETICS INC.
Query Watch 10.1%; Score 27.6; DB 4; Length 801;
Best Local Similarity 22.6%; Pred. No. 3.1e+02;
RESULT 1218
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.1%; Score 27.6; DB 6; Length 474; 51.9%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 516;
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(ZYMO ) ZYMOGENETICS INC.
(EYY Match 10.1%; Score 27.6; DB 4; Length 891;
(ery Match 22.6%; Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 888;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAC64074 standard; cDNA; 972 BP.
Mouse ion channel protein KCNE3 (MiRP2) cDNA, SEQ ID NO:7.
WO200063434-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human soft tissue sarcoma-upregulated DNA - SEQ ID 3496
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ID 7472
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Degenerate cDNA sequence for human UMLR variant #1
                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.1%; Score 27.6; DB 9; 56.7%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PD 23-AUG-2001.

PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

Query Match 10.1%; Score 27.6; DB 5;

Best Local Similarity 53.8%; Pred. No. 2.6e+02;

RESULT 1216
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AD024652 standard; DNA, 584 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ
W02004048938-A2.
10-1701-2004.
(PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AASO5961 standard; cDNA; 801 BP.
Degenerate cDNA sequence for human UMLR variant WC200130850-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 27.6; DB 4;
Pred. No. 3.2e+02;
                55.1%; Pred. No. 2.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABV56711 standard; cDNA; 516 BP.
Human prostate expression marker cDNA 56702.
WO200160860-A2.
                                                                                           ACH28744 standard; cDNA; 472 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADQ20676 standard; DNA; 481 BP.
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                                                                                                                                Human adult ovary cDNA #7124
US2003073623-A1.
                                                                                                                                                                                                       17-APR-2003.

(DRMA/) DRMANAC R T.

(LABA/) LABAT I.

(STAC/) STACHE-CRAIN B.

(DICK/) DICKSON M.C.

(JONE/) JONES L W.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 J8-A2.

2004.

2004.

EVATOR OF THE PROTEIN DES J8-Y MATCH BEST LOCAL SIMILATITY BEST LOCAL SIMILATITY BEST LOCAL SIMILATITY BEST LOCAL STANDS FOR WOOZOOJE FOR W
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Best Local Similarity
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Best Local Similarity RESULT 1212
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AAF71473 standard; DNA; 1575 BP.
Corynebacterium glutamicum SMP protein nucleotide sequence SEQ ID NO:227.
WO200100844-A2.
                                                                                             Proliferative glomerular nephritis-associated gene sequence SEQ ID:88.
WO200173022-A1.
04-OCT-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AALGE038 standard; cDNA; 1531 BP.

Human cell adhesion and extracellular matrix protein (CADECM)-26 cDNA.
WO200347526-A2.
12-JUN-2003.
(INCY-) INCYTE GENOMICS INC.
ery Match
10.1%; Score 27.6; DB 9; Length 1531;
st Local Similarity 51.6%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                       JT 1223
MD92292 standard; CDNA; 1044 BP.
Mouse hair keratin-associated-protein encoding cDNA SEQ ID NO:151
WO2003042387-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.1%; Score 27.6; DB 10; Length 1044; 51.6%; Pred. No. 3.4e+02;
                                                                                                                                                                       Luery Match
10.1%; Score 27.6; DB 4; Length 1014;
Beet Local Similarity 50.4%; Pred. No. 3.3e+02;
RESULT 1222
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                LOCAL MATCH 10.1%; Score 27.6; DB 2; Length 1315; Best Local Similarity 56.7%; Pred. No. 3.7e+02; RESULT 1226
                                                                                                                                                                                                                                                                                                                                                                                            Length 1034;
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Score 27.6; DB 3; Length 972;
Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAS90615 standard; cDNA; 1192 BP.
DNA encoding novel human diagnostic protein #26419.
WO200175067-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADA29218 standard; DNA; 1338 BP.
DNA encoding Acinetobacter baumannii protein #505
US6562958-B1.
                                                                                                                                                                                                                                                             AA74251 standard; cDNA; 1034 BP.

Human secreted protein gene 29 SEQ ID NO:39.

W0200056754-A1.

W22 SEP-2000.

(HUMAN GENOME SCI INC.

ery Match

10.1%; Score 27.6; DB 3;

st Local Similarity 51.6%; Pred. No. 3.46+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.1%; Score 27.6; DB 5; 49.3%; Pred. No. 3.5e+02;
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(GENO-) GENOME THERAPEUTICS CORP.
(ery Match
10.1%; Score 27.6; DB 9;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity 49.3%; Pred. No. 3.5e4
RESULT 1225
ID AAZA1312 standard; cDNA; 1315 BP.
DB Human normal ovarian tissue derived cDNA 91.
PN DE19816395-A1.
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Human peptidase, HPEP-17 coding sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       07-OCT-1999.
(META-) METAGEN GES GENOMFORSCHUNG MBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               22-MAY 2003.
(UYKE-) UNIV KEIO.
(NIPR-) JAPAN SOC PROMOTION SCI.
                                                                         ABA77081 standard; DNA; 1014 BP
10.1%;
50.8%;
                                                                                                                                                                    (KYOW ) KYOWA HAKKO KOGYO KK.
                 Best Local Similarity
RESULT 1221
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RESULT 1228
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RESULT 1229
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(HYSE-) HYSEQ INC.
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(BADI ) BASF AG.
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sednence

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Length 4459
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                                                                                             Length 4459;
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                                                                                                                                                                                                                                                                                                 AAC75103 standard, cDNA, 5245 BP.
Human ORFX ORF658 polynucleotide sequence SEQ ID NO:1315
WO200058473-A2.
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                                                                                                                                                                Human collapsin response mediator protein 2 gene 3' W02003040320-A2.
[15-MAY-2003.
[151S-] ISIS PHARM INC.
10.1%; Score 27.6; DB 10; Let Local Similarity 56.7%; Pred. No. 5.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABL70623 standard; DNA; 6045 BP. Chemically treated cell signalling DNA sequence#257 WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Human immune Bystem associated gene SEQ ID NO: 155 WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                             ABL32151 standard; DNA; 5276 BP.
Human immune system associated gene SEQ ID NO: 124
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABK31540 standard; DNA; 6045 BP. Signal transduction associated gene modified DNA W020020926-A2.
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(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 10.1%; Score 27.6; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   03-JAN-20UL.
(EPIG-) EPIGENOMICS AG.
2ry Match 10.1%; Score 27.6; DB 6;
10.1%; Score 27.6; DB 6;
                                                                                             Score 27.6; DB 9;
Pred. No. 5.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 27.6; DB 6;
Pred. No. 6.1e+02;
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Pred. No. 6.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                         Score 27.6; DB 3;
Pred. No. 6.1e+02;
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Pred. No. 6.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human polynucleotide SEQ ID NO 2384. 09-ARG-200157190-A2.
PD 19-FEB-2003.

PA (WARN ) WARNER LAMBERT CO.

QUETY MAtch
Best Local Similarity 56.7%; Pred.
RESULT 1239

ID ADC66333 standard; DNA; 4459 BP.
DE Human collapsin response mediator
PN WO2003040320-A2.
PD 15-MAY-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human polynucleotide SEQ ID NO 416. WO200157190-A2.
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ID AA199595 standard; DNA; 5690 BP.
DE Human polynuclectide SEQ ID NO 54
PN WO200155173-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DNA; 5763 BP.
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55.1%;
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51.6%;
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(EPIG-) EPIGENOMICS AG.
(YMATCh 10.1%;
) Tring timilarity 53.8%;
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Local Similarity 49.3%;
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ID ABL32182 standard; D: Human immune system ip WO200200928-A2.
PD 03-JAN-2002.
PA (EPIG-) EPIGENOMICS 2
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RESULT 1246
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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(HYSE-) HYSEQ INC.
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(HYSE-) HYSEQ INC.
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                                                                                                                                                                                                                                                                                                                                     SEQ ID
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAH54212 standard; DNA; 3043 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:3576
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAHS4954 standard; DNA; 3417 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:4318
WO200134809-A2.
                                                                                                                                                                                                                                                                2809;
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                                                                                                                 Length 1731;
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                                                                                                                                                                                                                                                                                                                                                                                                              Length 3039
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Human nervous system related polynucleotide SEQ ID NO 7922.
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                                                                                                                                                                                                                                                                                                                AAF21851 standard; DNA; 3039 BP.
Human breast and ovarian cancer associated antigen gene
Human breast and ovarian cancer associated antigen gene
21-0200055173-A1.
21-052-2000.
HUMAN GENOME SCI INC.
PLY MATCH
10.1%; Score 27.6; DB 3; Length
5t Local Similarity 56.7%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                Length
                                                                                                                                      Human ovarian tumor cDNA; 2809 BP.
DE19817557-A1.
21-OCT-1999.
(META-) METAGEN GES GENOMFORGUMENT.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABT13431 standard; DNA; 3361 BP.
Breast specific related polynucleotide SEQ ID No 146.
WO20027723-A2.
                                                                                                                 10.1%; Score 27.6; DB 3; 51.6%; Pred. No. 4.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           LTD.
10.1%; Score 27.6; DB 4;
50.8%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.1%; Score 27.6; DB 4; 50.8%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.1%; Score 27.6; DB 4; 72.0%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 27.6; DB 4;
Pred. No. 5.1e+02;
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Pred. No. 5.2e+02;
                                                                                                                                                                                                                                             GENOMFORSCHUNG MBH.
10.1%; Score 27.6; DB 2;
56.2%; Pred. No. 4.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.1%; Score 27.6; DB 5; 53.8%; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACF25352 standard; DNA; 4459 BP.
Human dihydropyrimidinase-related protein gene.
EP1284298-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
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17-MAY-ZOO1.
(GLAX ) GLAXO GROUP LTD.
(GLAX ) GLAXO SOUP LTD.
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52.6%;
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(JANC ) JANSEN PHARM NV.
                                                                            20-JUL-2000.
(INCY-) INCYTE PHARM INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        LTD.
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(DIAD-) DIADEXUS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         17-MAY-2001.
(GLAX ) GLAXO GROUP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   17-MAY-2001.
(GLAX ) GLAXO GROUP
                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 1231
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Best Local Similarity
RESULT 1234
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Best Local Similarity
RESULT 1235
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RESULT 1236
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RESULT 1230
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RESULT 1233
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                                                             WO200042201-A2.
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Best Local Si
RESULT 1238
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Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36319.
WO200157182-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Signal transduction associated gene modified complementary DNA #13. WO200200926-A2. 03-JAN-2002. (BPIG-) EPIGENOMICS AG.
       E Human lymphoid cell proliferative disorder gene derived DNA #133.

N WO2003034226-A2.

D 30 NAY-2003.

A 3DFIGEO BEIGENOMICS AG.

Query Match

Best Local Similarity 52.6%; Pred. No. 7.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 27.6; DB 6; Length 15698; Pred. No. 9e+02;
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(EPIG-) EPIGENOMICS AG.
10.1%; Score 27.6; DB 6; Length 15881;
r ronal Similarity 53.8%; Pred. No. 9e+02;
                                                                                                                                                                                                                                                                                                                                                                                        Length 10594;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.1%; Score 27.6; DB 3; Length 12494; 53.8%; Pred. No. 8.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.1%; Score 27.6; DB 6; Length 15881; 53.8%; Pred. No. 9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 10598;
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                                                                                                                                                                                                                                       Length 9465;
                                                                                                                                                                                                                                                                                        AAZ49991 standard; DNA; 10594 BP.
Plasmid pGN205 for inhibition of unc-22 expression in C. WO200001846-A2.
13-JAN-2000.
(DEVG-) DEVGEN NV.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Plasmid pCN110 for expression of T7 RNA polymerase in C. MO200001846-A2.
13-CAN-2000.
ic Ebcy-) DEVGEN NV.
                                                                                                                                                                                                                                                                                                                                                                                                                                             DNA; 10598 BP. inhibition of unc-22 expression in C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         2114
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             MCACCIO.

09-AUG-2001.

(HUMA.) HUMAN GENOME SCI INC.

(HIMA.) HUMAN GENOME SCI INC.

10.1%; Score 27.6; DB 4;

(ery Match

cimilarity 52.6%; Pred. No. 8.1e+02;
                                                                                                                                                                                                                                       Score 27.6; DB 3;
Pred. No. 7.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                        10.1%; Score 27.6; DB 3; 53.8%; Pred. No. 7.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human immune system associated gene SEQ ID NO:
                                                                                                                                         AAAS1008 standard; DNA; 9465 BP.
Simian immunodeficiency virus SIVrcm.
WO200034529-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA; 15881 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DNA; 15698 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DNA; 15881 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAK81507 standard; DNA; 11718 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DEPLOY EPIGENOMICS AG.

10.1%;
17 Match 10.1%;
17 Incal Similarity 50.0%;
                                                                                                                                                                                                                                       10.1%;
                                                                                                                                                                                                                                                      46.4%;
                                                                                                                                                                                                  15-JUN-2000.
(UABR-) UAB RES FOUND.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                Plasmid pGN207 for i. WO2000001846-A2. 13-JAN-2000. (DEVG-) DEVGEN NV.
                                                                                                  Best Local Similarity RESULT 1257
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity
RESULT 1264
ID ABK31183 standard; I
                                                                                                                                                                                                                                                     Local Similarity
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RESULT 1259
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RESULT 1262
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RESULT 1263
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ID AAZ49988 standard;
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                                                                                                                                                                                                                                                                                                                                                                                                                                             AAZ49992 standard;
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                                                                                                                                                                                                                                       Query Match
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Chemically pretreated complementary DNA associated with cell cycle #76.
WO200168911-A2.
                                         Best Local Similarity 53.8%; Fred. No. 6.46+02; RESULT 1248
ID ABZ10.05 standard; DNA; 6289 BP.
DB Haematopoietic cell proliferation disorder related DNA sequence #345.
                                                                                                                                                                                                                                                                                                                                                                                                      Score 27.6; DB 12; Length 6539;
Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 6543;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.1%; Score 27.6; DB 6; Length 6876; 55.1%; Pred. No. 6.7e+02;
                         Length 6045;
                                                                                                                                 (BPIG-2002.
(BPIG-) EPIGENOMICS AG.
(EPY Match 10.1%; Score 27.6; DB 8; Length 6289;
(ET) Match 10.1%; Pred. No. 6.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 27.6; DB 6; Length 7934; Pred. No. 7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 7143;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DNA transcription associated complementary genomic DNA #85.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Human gene regulation-associated gene oligonucleotide #299.
WO200177375-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABL33764 standard; DNA; 6876 BP.
Human immune system associated gene SEQ ID NO: 1737.
WC200200928-A2.
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Human immune system associated gene SEQ ID NO: 955-
WO200200928-A2.
                         Score 27.6; DB 6;
Pred. No. 6.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.1%; Score 27.6; DB 6; 56.7%; Pred. No. 6.6e+02;
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No. 6.8e+02;
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Pred. No. 7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 27.6; DB
Pred. No. 7e+02;
                                                                                                                                                                                                                                                      Mouse lymphoma associated, LA, contig #4.
US2003224460-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Lymphona associated polynucleotide #131.
WO200224867-A2.
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Pred.
                                                                                                                                                                                                                                   ADM79326 standard; cDNA; 6539 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADE84197 standard; DNA; 9289 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                DNA; 6543 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAS61344 standard; DNA; 7934 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              18-OCT-ZUUL.
(EPIG-) EPIGENOMICS AG.
ry Match 10.1%;
(EPIG-) EPIGENOMICS AG.
2TV Match 10.1%;
at Local Similarity 53.8%;
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56.7%;
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20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
10.1%;
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51.6%;
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(PEDE/) PEDERSEN F S.
(SORE/) SORENSEN A B.
(HERN/) HERNANDEZ J M.
(NIEL/) NIELSEN A A.
(MOVI/) MOVING H.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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(EPIG-) EPIGENOMICS AG.
                                                                                            Jieta

7272-A2.

T-2002.

JE-2002.

Y MATCH

SULF 1249.

ADM79326 standard.

DE Mouse 1ymphoma 7.

PN US203224460-7.

PN US203224460-7.

PN (FEDE/)

PA (FEDE/)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (EPIG-) EPIGENOMICS AG.
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(UYAA-) UNIV AARHUS.
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Best Local Similarity
RESULT 1255
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RESULT 1256
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RESULT 1250
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RESULT 1251
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RESULT 1253
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1D ABL32982
DE Human in
PN WO200200
PD 03-JAN-2
PA (BPIG-)
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elegans

elegans

elegans

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Human multidrug resistance associated protein gene associated sequence.
WO200257410-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.1%;
52.6%;
                              PD 25-JUL-2002.

PA (DNAS-) DNA SCI LAB INC.

QUETY MATCh 10.1%;

Best Local Similarity 52.6%;

RESULT 1275
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.1%;
55.1%;
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50.0%;
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58.5%;
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55.1%;
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55.1%;
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50.0%;
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48.1%;
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55.1%;
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Human MDR1 related
WO2003013537-A2.
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Best Local Similarity
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Best Local Similarity
RESULT 1282
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Best Local Similarity
RESULT 1288
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Best Local Similarity
                                                                                                                                                                                                              RESULT 1276
ID ADB20860 standard;
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.1%; Score 27.6; DB 9; Length 96596; 46.8%; Pred. No. 1.7e+03;
                                                                                                                                                    AG.
10.1%; Score 27.6; DB 6; Length 15881;
53.8%; Pred. No. 9e+02;
                                                                                                                                                                                                                                                                                      Length 19659;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 33146;
                                                 Length 15881;
                                                                                                                                                                                                                                                                                                                                                                                                         Length 20579;
                                                        Human gene regulation-associated gene oligonucleotide #25.
18-OCT-2001.
18 CT-2001.
17 Match
                                                                                                                                                                                                                                                                                                                              ABQ67073 standard; DNA; 20579 BP.
Human angiogenesis associated polynucleotide SEQ ID NO 103
WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADA03068 standard; DNA; 96595 BP.
Human PPP3CC carcinoma associated gene, SEQ ID NO:1586
WO2003057146-A2.
17-UUL-2003.
                                                                                                                                                                                                                            740.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Score 27.6; DB 11;
Pred. No. 1.7e+03;
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Pred. No. 1.7e+03;
                                                 10.1%; Score 27.6; DB 6; 53.8%; Pred. No. 9e+02;
                                                                                                                                                                                                                                                                                                                                                                      13.JUN-2002.
(BPIG-) EPIGENOMICS AG.
ery Match
10.1%; Score 27.6; DB 6;
ery match
10.1%; Pred. No. 9.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.1%; Score 27.6; DB 6; 48.1%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                      Score 27.6; DB 6;
Pred. No. 9.8e+02;
                                                                                                                                                                                                           ABL32767 standard; DNA; 19659 BP.
Human immune system associated gene SEQ ID NO:
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                  RESULT 1269

ID ABQ67191 standard; DNA; 33146 BP.

DE Listeria innocua contig DNA sequence #4.

PD IJSTER 170228891-A2.

PD 11-APR-2002.
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Human PPP3CC gene genomic DNA seguence.
WO2003053224-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADL27146 standard; DNA; 96593 BP. Human genomic sequence for PPP3CC US2003216558-A1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA; 96595 BP
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(CNRS ) CNRS CENT NAT RECH SCI.
                                                                                                                                                                                                                                               WOZUGECT.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.1%;
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46.8%;
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46.8%;
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(SAGR-) SAGRES DISCOVERY.
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(SAGR-) SAGRES DISCOVERY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (SAGR-) SAGRES DISCOVERY.
                  10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           (MORR/) MORRIS D W. (ENGE/) ENGELHARD E K.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADB72806 standard;
Human PPP3CC gene.
WO2003008583-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 1273
                                                                                                                                                                              Best Local Similarity RESULT 1267
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RESULT 1270
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RESULT 1271
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                                                          Best Local Similarity RESULT 1266
                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 1268
      WO200202807-A2.
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PD
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PD
PA
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ADB87949 standard; DNA; 98472 BP.
Human UGTIAL gene sequence SEQ ID NO:673.
20-FBB-2003.
(ERID-) EPIDAUROS BIOTECHNOLOGIE AG.
BY MATCH
10.1%; Score 27.6; DB 10; Length 98472; St Local Similarity 52.6%; Pred. No. 1.7e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 110000;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 98472;
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Length 98472
                                                                    NO:673
                                               ACF62745 standard; DNA; 98472 BP.
Cancer based on CYP3A5 related polynucleotide SEQ ID
W0203013534-A2.
20-FBE-2003.
(BPID-) BPIDAUROS BIOTECHNOLOGIE AG.
EPT MAtch
10.1%; Score 27.6; DB 8; Leng
st Local Similarity 52.6%; Pred. No. 1.7e+03;
                                                                                                                                                                      AUBZUBEO standard; DNA; 98472 BP.
MRP1 based cancer related nucleic acid SEQ ID NO:673.
W0.201301333-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   QO-FEB-2003.
(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
ery Match
10.1%; Score 27.6; DB 10;
ery Match
52.6%; Pred. No. 1.7e+03;
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No. 1.8e+03;
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(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
Query Match
10.1%; Score 27.6; DB 8;
Best Local Similarity 52.6%; Pred. No. 1.7e+03;
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No. 1.8e+03;
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No. 1.8e+03;
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No. 1.8e+03;
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No. 1.8e+03;
Score 27.6; DB 6;
Pred. No. 1.7e+03;
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Pred. No. 1.8e+03;
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No. 1.8e+03;
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No. 1.8e+03;
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No. 1.8e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADB92123 standard; DNA; 98472 BP.
Human MDR1 related DNA sequence SEQ ID NO:673.
WO2003013535-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DNA; 98472 BP.
DNA sequence SEQ ID NO:673
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    20-FEB-2003.
(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
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Pred.
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Pred.
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Pred.
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Pred.
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10.1%; Score 27.6; DB 10; Length 128993; 52.6%; Pred. No. 1.9e+03;
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                Length 110000;
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(BPID-) BPIDAUROS BIOTECHNOLOGIE AG.
(ery Match 10.1%; Score 27.6; DB 8; Length 128993;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 115218;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ACF62749 standard; DNA; 128993 BP.
Cancer based on CYP3A5 related polynucleotide SEQ ID NO:681.
WC2003013534-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                MRP1 based cancer related nucleic acid SEQ ID NO:681.
WO2003013533-A2.
20-FEB-2003.
                                                                                                                                                                             Score 27.6; DB 12;
Pred. No. 1.8e+03;
                                                                                                                                                                                                                                                                                                                                       Score 27.6; DB 12;
Pred. No. 1.8e+03;
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Human MDR1 related DNA sequence SEQ ID NO:681.

MO2003013537-A2.

20-PEB-2003.

(EPID-)

EPIDAUROS BIOTECHNOLOGIE AG.

(EPID-)

ery Mac.

10.1%; Score 27.6; DB 10;

st Local Similarity 52.6%; Pred. No. 1.9e+03;
                Score 27.6; DB 12;
Pred. No. 1.8e+03;
                                                                    Score 27.6; DB 12;
Pred. No. 1.8e+03;
                                                                                                                                                                                                                               Score 27.6; DB 12;
Pred. No. 1.8e+03;
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Pred. No. 1.8e+03;
                                                                                                                          27.6; DB 12;
No. 1.8e+03;
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Pred. No. 1.8e+03;
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BAC containing repeats from centromeres 1-4 #19.
WC200055325-A2.
                                                                                                                                                                                                                                                                                                                                                                                        ACA64845 standard; DNA; 115218 BP.
Human HNRNP GP43 DNA corresponding to AL034397.
DE10127572-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADB92131 standard; DNA; 128993 BP.
Human MDR1 related DNA sequence SEQ ID NO:681.
WO203031355-A2.
30-FEB-2003.
(BPID-) EPIDAUROS BIOTECHNOLOGIE AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADBB7957 standard; DNA; 128993 BP.
Human UGT1A1 gene sequence SEQ ID NO:681.
WO2003013536-A2.
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(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
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Pred.
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50.0%;
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50.0%;
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55.1%;
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50.0%;
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50.0%;
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(PATH-) PATHOARRAY GMBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
Best Local Similarity
RESULT 1297
ID ACR62749 standard; DN
DE Cancer based on CYP3A
PN WC2003115334-A2.
PD 20-FEB-2003.
PA (EPID-) EPIDAUROS BIO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1298
ID ADB20868 standard; DN
DE MRP1 based cancer rel
PN WC2003013533-A2.
PD 20-FEB-2003.
PA (EPID-) EPIDAUROS BIO
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 1299
ID ADB87957 standard; DN
DE Human UGT1A1 gene seq
PN WO2003013536-A2.
PD 20-FEB-2003.
PA (EPID-) EPIDAUROS BIOT
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RESULT 1300
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Best Local Similarity
RESULT 1295
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Best Local Similarity
RESULT 1296
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Best Local Similarity
RESULT 1292
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Best Local Similarity
RESULT 1293
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Best Local Similarity
RESULT 1294
                Query Match
Best Local Similarity
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Best Local Similarity
RESULT 1291
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ESULT 1289
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Length 183178;
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                                                Length 129021;
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Human soft Lissue sarcoma-upregulated DNA - SEQ ID 1263
W02004048938-A2.
10-JUN-2004.
[PROT-) PROTEIN DESIGN LABS INC.
ET MATCH.
10.0$; Score 27.4; DB 12; Length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAH68533 standard, DNA, 349980 BP.
C glutamicum coding sequence fragment SEQ ID NO: 7068.
EP1108790-A2.
                                                                                                                ADL13873 standard; DNA; 183178 BP. Osteoarthritis-associated polymorphic nucleotide #405.
                                                                                                                                                                                                                                                                                                                                                                         02-JAN-2003.
(NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
(ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
PRY MATCH 10.1%; Score 27.6; DB 10.
It Local Similarity 53.8%; Pred. No. 2.5e+03.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PA (NAAD-) NAT INST ADVANCED IND SCI & TECHNOLOGY.
PA (ADSC-) CENT ADVANCED SCI & TECHNOLOGY INCUBATIO.
QUERY MAtch 10.1%; Score 27.6; DB 10;
Best Local Similarity 53.8%; Pred. No. 2.5e+03;
RESULT 1306
                                                                                                                                                                                                          (INCY-) INCYTE GENOMICS INC.
10.1%; Score 27.6; DB 10;
it Local Similarity 51.6%; Pred. No. 2.1e+03;
                                              Query Match 10.1%; Score 27.6; DB 3;
Best Local Similarity 52.6%; Pred. No. 1.9e+03;
RESULT 1303
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADC87621 standard; DNA; 349938 BP.
Human GPCR related polynucleotide SEQ ID NO:2074.
EP1270724-A2.
02-JAN-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             20--UN-2001.
(KYOW ) KYOWA HAKKO KOGYO KK.
(ETY MAtch 10.1%; Score 27.6; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.4; DB 10; 50.4%; Pred. No. 2.2e+02;
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
10.0%; Score 27.4; DB 5;
ref foral Similarity 43.7%; Pred. No. 2.8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.0%; Score 27.4; DB 5; 43.7%; Pred. No. 2.8e+02;
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ID ADL37205 standard; DNA; 433 BP.
DE Human ovarian cancer DNA marker #11095.
PN W0200170979-A2.
PD 27-SEP-2001.
PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADH84487 standard; DNA; 201 BP.
Enterococcus faecalis polynucleotide #2372.
US6517156-B1.
09-SEP-2003.
(DOUC/) DOUCETTE-STAMM L A.
(BUSH/) BUSH D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human ovarian cancer DNA marker #4798.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                       ADC86940 standard; DNA; 349901 BP.
Human GPCR gene SEQ ID NO:1393.
EP1270724-A2.
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21-SEP-2000.
(UYCH-) UNIV CHICAGO.
                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 1304
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 1307
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03-JUL-2003.
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Best Local Si
RESULT 1308
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Length

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10.0%;
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(SCAN/) SCANLAN M J.
(LEES/) LEE S.
(OLDL/) OLD L J.
Query Match
Best Local Similarity
RESULT 1320
ID ABT11573 standard, DI
DE Yeast selected inter:
PN WQ200266504-A2.
PD 29-AUG-2002.
PA (HYBR-) HYBRIGENICS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAS85335 standard; c
DNA encoding novel h
WO200175067-A2.
11-OCT-2001.
                                                                                                        29-AUG-2002. (HYBR-) HYBRIGENICS.
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(HYBR-) HYBRIGENICS.
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RESULT 1322
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RESULT 1324
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Best Local Similarity
RESULT 1325
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RESULT 1326
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                                                                                                                                         Query Match
Best Local Similarity
RESULT 1321
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACA24780 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                      WO200229113-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               06-JAN-2004
                                                                                                                                                                                                                                                                                   Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABQ41296 standard; DNA; 617 BP.
Oligonucleotide for detecting cytosine methylation SEQ ID NO 27887.
WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABQ41297 standard; DNA; 617 BP.
Oligonucleotide for detecting cytosine methylation SEQ ID NO 27888.
WO200218632-A2.
Human immune/haematopoietic antigen encoding cDNA SEQ ID NO:7116. W0200157182-A2. O9-AUG-2001. (HUMA) HUMAN GENOME SCI INC. (HUMA) HUMAN GENOME SCI INC. 27.4; DB 4; Length 457; Et Local Similarity 57.6%; Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              SEQ ID NO:185
                                                                                                                                                                                                                                                                                                                                                                                   Length 462;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 468;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 559
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                                                                                                                                                                                                                                                                                                                                                         ID 5761
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAC53727 standard; DNA; 559 BP.
AAC53727 standard; DNA; 559 BP.
EPI0334055.82.
06-SEP-2000.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAC93690 standard; cDNA; 498 BP. Cat flea hindgut and Malpighian tubule (HMT) cDNA, WO200061621-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 MCZCCZO.

(RPTG-) BPIGENOMICS AG.

(EPTG-) BPIGENOMICS AG.

10.0%; Score 27.4; DB 6;

ery Match

10.0%; Score 27.4; DB 6;

ery Match

10.0%; Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                 10.0%; Score 27.4; DB 9; 62.3%; Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MCACCL.
25-JUL 2002.
(BICC-) BIOCARDIA INC.
(ery Match 10.0%; Score 27.4; DB 6;
(ery Match 50.0%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.0%; Score 27.4; DB 3; 65.6%; Pred. No. 3e+02;
                                                                                                                                                                                                                                                                                                                  RESULT 1313
ID ADQ22941 standard; DNA; 466 BP.
ID Human soft tissue sarcoma-upregulated DNA - SEQ
PN WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%; Score 27.4; DB 5; 55.9%; Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.4; DB 3; 57.6%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human leukocyte derived cDNA, 530 BP.

WOZOS7414-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAS21889 standard; DNA; 468 BP.
Human collagen gene COL9Al intron 10.
US6265157-Bl.
24-UUL-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (UYAL-) UNIV ALLEGHENY HEALTH SCI. (UYJE-) UNIV JEFFERSON THOMAS. (UYOU-) UNIV OULU.
                                                                                                                           ACH18968 standard; cDNA; 462 BP.
Human adult heart cDNA #3282.
US2003073623-A1.
                                                                                                                                                                          PD 17-APR-2003.
PA (DRMA/) DRMANAC R T.
PA (LABA/) LABAT I.
PA (STAC/) STACHE-CRAIN B.
PA (DICK/) DICKSON M C.
PA (JONE/) JONES L W. 10.0C
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                07-MAR-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                    Best Local Similarity RESULT 1312
                                                                                                                                                                                                                                                                                                                                                                                                                                          Best Local Similarity RESULT 1314
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             19-OCT-2000.
(HESK-) HESKA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
RESULT 1317
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RESULT 1319
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645.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.0%; Score 27.4; DB 12; Length 935; 55.9%; Pred. No. 3.8e+02;
                                                         ABTI1573 standard; DNA; 634 BP.
Yeast selected interacting domain coding sequence SEQ ID NO:
WO200266504-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 1002;
                                                                                                                                                                                                                                                                                                                                        Length 635;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAL51357 standard; DNA; 1000 BP.
Haemophilus influenzae BASB229 gene-related DNA sequence
WO2002100891-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADL14028 standard; cDNA; 935 BP.
Human cDNA encoding sarcoma-associated antigen NY-SAR-27
US2004063101-A1.
                                                                                                                                                                      Length
                                                                                                                                                                                                                                                                                                                                                                                                    ABK73027 standard; DNA; 650 BP.
Bacillus licheniformis genomic sequence tag (GST) #318
                                                                                                                                                                                                                                ABS62968 standard; DNA; 635 BP.
Selected Interacting Domain (SID) polynucleotide #165.
WO200259255-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         cDNA; 1002 BP.
human diagnostic protein #21139
10.0%; Score 27.4; DB 6; 52.1%; Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                        10.0%; Score 27.4; DB 6; 75.6%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.0%; Score 27.4; DB 5; 53.2%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 27.4; DB 8;
Pred. No. 3.8e+02;
                                                                                                                                                                    Score 27.4; DB 6;
Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.0%; Score 27.4; DB 6; 54.5%; Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADL02866 standard; DNA; 1080 BP.
DNA encoding a M. catarrhalis protein #552.
US6673910-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (GENO-) GENOME THERAPEUTICS CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Prokaryotic essential gene #6437 WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DNA; 1086 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (NOVO ) NOVOZYMES BIOTECH INC. (NOVO ) NOVOZYMES AS.
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03-007-2002.
(ELITEA PHARM INC.
""MATCh 10.0%;
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(HUMA-) HUMAN GENOME SCI INC.
ry Match
t Local Similarity 62.3%; Pred. No. 4.5e+02;
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RESULT 1346
ID ADQ99420 standard; CI
DE DNA encoding human GF
PN US6569662-B1.
PD 27-MAY-2003.
PA (HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            27-AUG-2003.
(CELL-) CELLZOME AG.
                                Query Match
Best Local Similarity
RESULT 1338
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity
RESULT 1343
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RESULT 1344
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RESULT 1345
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(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                    Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                 ABZ22252 standard; cDNA; 1086 BP. Nucleotide sequence GI4885400 related to holocytochrome C synthase. WO200299054-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AD22121 Standard; CDNA; 1086 BP.
Nucleotide sequence GI1209634 related to holocytochrome C synthase.
WO200299054-A2.
                                                                                                                                                   ABK84102 standard; cDNA; 1086 BP.
Human cDNA differentially expressed in granulocytic cells #873
WO200228999-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10-JUN-2004.

10-JUN-2004.

(PROT-) PROTBIN DESIGN LABS INC.

(PROT-) PROTBIN DESIGN LABS INC.

(Ery Match

(Ery M
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                                                             10.0%; Score 27.4; DB 6; Length 1086; 54.5%; Pred. No. 4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.0%; Score 27.4; DB 4; Length 1521; 45.0%; Pred. No. 4.5e+02;
                                                                                                                                                                                                                                                                                                              Score 27.4; DB 6; Length 1086; Pred. No. 4e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 1086
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WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAL99442 standard; DNA; 1521 BP.
Human excretory related polynucleotide SEQ ID NO 1206.
WOZO0155313-A2.
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Human bladder related polynucleotide, SEQ ID NO: 120.
WO200159064-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              #
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WO200055350-A1.
21-SEP-2000.
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Human bladder associated antigen #16 genomic DNA
US2003199008-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     y Match 10.0%; Score 27.4; DB 8; Local Similarity 54.5%; Pred. No. 4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%; Score 27.4; DB 4; 45.0%; Pred. No. 4.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.0%; Score 27.4; DB 8; 54.5%; Pred. No. 4e+02;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAT68079 standard; DNA; 1200 BP.
H. pylori cytoplasmic protein ORF 11ep12011orf9.
WO9640893-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAC78091 standard; cDNA; 1566 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADO19774 standard; DNA; 1086 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  RESULT 1331

ID ABZ23251 standard; cDNA; 1086

DE Nucleotide sequence G11209634

PD 12-DEC-2002.

PA (EXEL-) EXELIXIS INC.
                                                                                                                                                                                                                          WOZUCZ-.
11-APR-2002.
(GENE-) GENE LOGIC INC.
Orv Match 10.0%;
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(HUMA-) HUMAN GENOME SCI INC.
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(HUMA-) HUMAN GENOME SCI INC.

Query Match 10.0%;
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(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         12-DEC-2002.
(EXEL-) EXELIXIS INC.
13-DEC-2001.
(AVAL-) AVALON PHARM.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (EXEL-) EXELIXIS INC.
                                                         Query Match
Best Local Similarity
RESULT 1329
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1332
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Best Local Similarity
RESULT 1336
                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
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Best Local Similarity
RESULT 1333
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RESULT 1334
ID AA199442 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
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A (ASTR ) ASTRA AB.
Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     12-DEC-2002
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                                                                                                                                                                                                                                                                                                                                                                 RESULT 1330
ID ABZ2325;
DE Nucleot:
PN WO200299
PD 12-DEC-;
PA (EXEL-)
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Length 1833;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.0%; Score 27.4; DB 10; Length 2000; 59.7%; Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 10; Length 2000; 59.7%; Pred. No. 4.9e+02;
Length 1566;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 1759;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.0%; Score 27.4; DB 2; Length 2009; 55.9%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.0%; Score 27.4; DB 4; Length 2030; 55.9%; Pred. No. 5e+02;
                                                                                                                                                              Length 1566;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AD23310 standard; DNA; 1833 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 6730.
#P02004048938-A2.
10-MUN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
                                                         AAH33534 standard; cDNA; 1566 BP.
Human colon cancer antigen encoding cDNA SEQ ID NO:590.
WO200122920-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Disease treating protein complex-derived gene #1490 EP1338608-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      vuery Match 10.0%; Score 27.4; DB 12; Best Local Similarity 54.5%; Pred. No. 4.8e+02; RESULT 1342
                                                                                                                    05-APR-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 10.0%; Score 27.4; DB 4;
st Local Similarity 62.3%; Pred. No. 4.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                  PD 25-JUL-2002.

PA (INCY-) INCYTE GENOMICS INC.

Query March

Best Local Similarity 53.2%; Pred. No. 4.7e+02;

RESULT 134
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  cDNA; 2030 BP.
GPCR-like protein seqid 1090
                                                                                                                                                                                                                                                                                                                                                                                   ABS51362 standard; cDNA; 1759 BP. cDNA encoding human secretory protein #60.WO200257304-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  23-SEP-1999.
(META-) METAGEN GES GENOMFORSCHUNG MBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAZ33670 standard; cDNA; 2009 BP.
Human breast tumour-associated EST 60.
DE19813839-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   AAIS9197 standard; cDNA; 2030 BP.
Human polynucleotide SEQ ID NO 1400.
WO200153312-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ACC61838 standard, DNA; 2000 BP. Gene sequence #SEQ ID 2458.
EP1258494-A1.
20-NOV-2002.
CELL-) CELLZOME AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADK63771 standard; DNA; 2000 BP.
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Query Match Best Local Si RESULT 1348

Query Match

Ouery Match

Query Match

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10.0%; Score 27.4; DB 12; Length 3418; 50.4%; Pred. No. 6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 2804;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 10; Length 2790; 54.5%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 2790;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 3239;
                   Length 2790;
                                                                                                                                                                         Length 2790;
                                                                                                                                                                                                                                                                                                                             Length 2790;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 2848;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            RESULT 1360
ID AAC76844 standard; cDNA; 2848 BP.
DB Human ORFX ORP2399 polynuclectide sequence SEQ ID NO:4797
PN WO20005-8473-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human airway trypsin-like protease HAT coding sequence. WO2004053496-A1.
24-JUN-2004.
(HIRZL)/ HINZMANN B.
(HERM/) HEDBEN E.
                                                                                                                                                                                                                                                                                                                                                                                  ADD66387 standard; cDNA; 2790 BP.
Human lung tumour-specific related cDNA, SEQ ID No
WC200292001-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.4; DB 10; 54.5%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 27.4; DB 12;
Pred. No. 5.6e+02;
                                                                                                                                                                         10.0%; Score 27.4; DB 3; 54.5%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 3; 55.9%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                           10.0%; Score 27.4; DB 4; 54.5%; Pred. No. 5.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  10.0%; Score 27.4; DB 4; 55.9%; Pred. No. 5.9e+02;
                     10.0%; Score 27.4; DB 2; 54.5%; Pred. No. 5.6e+02;
                                                                                                                                                                       Best Local Similarity 54.5%; Pred. No. 5.6e+02; RESULT 1356
DAD23170 standard; cDNA, 2790 BP.
DE Human Lung tumour-specific protein L86S-36 cDNA. PN WC200172295-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAH14011 standard; cDNA; 3239 BP.
Human cDNA sequence SEQ ID NO:11103.
EP1074617-A2.
                                                                            AAC79095 standard; cDNA; 2790 BP.
Human lung tumour-specific cDNA #48.
WO200060077-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADE87641 standard; cDNA; 2790 BP.
Human lung tumour antigen cDNA #48.
US2003118599-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DNA; 3418 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADG31188 standard; DNA; 3418 BP Novel mouse gene #13. W02003089644-A1. 30-OCT-2003. (RIKE) RIKEN KK. (DNAF-) DNAFORM KK. (MITU) MITSUBISHI CHEM CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (HELI-) HELIX RES INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         0 05-OCT-2000.
(CURA-) CURAGEN CORP.
Query Match
     (CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                         21-NOV-2002.
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (HERM/) HERMANN K. (ROSE/) ROSENTHAL A.
                                                                                                                                     12-OCT-2000.
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                           (CORI-) CORIXA CORP
                 Query Match
Best Local Similarity
RESULT 1355
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 1358
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 1363
                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
RESULT 1357
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Local Similarity
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RESULT 1361
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 1359
ID ADQ30691 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                26-JUN-2003
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           07-FEB-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
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                                                                                                                                                                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 2358;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 27.4; DB 10; Length 2358; Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 2358;
     10.0%; Score 27.4; DB 5; Length 2030; 55.9%; Pred. No. 5e+02;
                                                                                                                                                                                                                                  10.0%; Score 27.4; DB 9; Length 2030; 55.9%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                           10.0%; Score 27.4; DB 6; Length 2065; 57.6%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 4; Length 2182; 62.3%; Pred. No. 5.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 2597;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity 62.3%; Pred. No. 5.4e+02; RESULT 1354
ID AAZ07171 standard; CDNA; 2790 BP.
DE Human Lung tumour protein L86S-36 extended CDNA sequence. PD 05-AUG-1999.
                                                                                                                                                                                                                                                                                       ABS64932 standard; cDNA; 2065 BP.
Soybean ribonuclease D-like (RNaseD-like) cDNA #3.
US2002088026-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 27.4; DB 10;
Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 27.4; DB 10;
Pred. No. 5.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity 57.6%; Pred. N
RESULT 1349

ID AAHIS656 standard; CDNA; 2182 BP.

DE Human CDNA sequence SEQ ID NO:14003.

PN EP1074617-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAH18621 standard; CDNA; 2597 BP.
Human CDNA Bequence SEQ ID NO:18837.
EP1074617-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADE56467 standard; DNA; 2358 BP.
Human gene D88674, SEQ ID NO 2320.
WO2003016475-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADE60753 standard; DNA; 2358 BP.
Human gene D88674, SEQ ID NO 6665.
WO2003016475-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADE60749 standard; DNA; 2358 BP.
Human gene D88674, SEQ ID NO 6661.
WO2003016475-A2.
                                               Novel human cDNA SEQ ID NO 1090. US2003104529-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.0%;
83.8%;
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83.8%;
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83.8%;
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PA (GEHO ) GEN HOSPITAL CORP.
PA (FARB ) BAYER AG.
Query Match
Best Local Similarity 83.8%
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         27-FEB-2003.
(GEHO ) GEN HOSPITAL CORP.
(FARB ) BAYER AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CATHERS 2003.

(GEHO ) GEN HOSPITAL CORP.

(FARB ) BAYER AG.
                                                                                                                                                                                                                                                                                                                                        04-JUL-2002.

04-JUL-2002.

(BUTL/) BUTLER K H.

(CAHO/) CAHOON R E.

(RAFA/) RAFALSKI J E.

(SAKA/) SAKAI H.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           07-FEB-2001.
(HELI-) HELIX RES INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           07-FEB-2001.
(HELI-) HELIX RES INST.
                                                                                                                                                                                                                DRMANAC R T.
Query Match
Best Local Similarity
RESULT 1347
                                                                                                                                 (ZHOU/) ZHOU P.
(TANG/) TANG Y T.
(LIUC/) LIU C.
(ASUN/) ASUNDI V.
(DRMA/) DRMANAC R T
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 1350
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
RESULT 1352
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
RESULT 1351
                                                                                                                                                                                                                                                    Local Similarity
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Length 4178;

Length 4342;

Length 4041;

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ABL30304 standard; DNA; 4472 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO 42385-
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Drosophila melanogaster genomic polynucleotide SEQ ID NO 31177.
WO200171042-A2.
27-SEP-2001.
     Score 27.4; DB 12; Length 3646; Pred. No. 6.1e+02;
                                                        AAH54209 standard; DNA; 4041 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:3573.
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                           S. epidermidis genomic polynucleotide sequence SEQ ID NO:3989. WO200134809-A2.
                                                                                                                                                         RESULT 1373

ID AD036310 standard, DNA, 4178 BP.

DE Intracellular antibody isolation-related KAN gene #82.

PD MO2004046192-A2.

PRESULT 1374, Pred. No. 6.4e+02;

RESULT 1374
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAI99444 standard; DNA; 4507 BP.
Human excretory related polynucleotide SEQ ID NO 1208.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AMIGAUBS standard; CDNA; 4507 BP.
Human bladder related polynucleotide, SBQ ID NO: 122.
WO200159064-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                PD 02-AUG-2001.
PA (HUMA-) HUMAN GENOME SCI INC.
Query Match 10.0%; Score 27.4; DB 4;
Best Local Similarity 45.0%; Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADF71687 standard; DNA; 4507 BP.
Human bladder associated antigen #16 genomic DNA
US2003199008-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DB 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 27.4; DB 4;
Pred. No. 6.5e+02;
                                                                                                          17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
(GLAX ) GLAXO GROUP LTD.
(ery Match 10.0%; Score 27.4; DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%; Score 27.4; DB 4; 46.6%; Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.0%; Score 27.4; DB 4; 52.1%; Pred. No. 7.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.0%; Score 27.4; DB 4; 45.0%; Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           23-0CT-2003.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) Autch 10.0%; Score 27.4; DB 12
lery Match 15.0%; Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADN95328 standard, DNA, 5572 BP.
Human BEC/LEC-related gene sequence SeqID250.
WO2003080640-Al.
                                                                                                                                                                                                                                                                                                                                                         AAH54625 standard; DNA; 4342 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                02-0CT-2003.
(LUDW-) LUDWIG INST CANCER RES.
(LICN ) LICENTIA LTD.
     10.0%;
50.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                              17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
Query Match
Best Local Similarity
RESULT 1372
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 1375
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
Best Local Similarity
RESULT 1378
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
RESULT 1379
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 1380
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     27-SEP-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
                                                                                                                                                                                                                                                                                         ABLI0838 standard; cDNA; 3583 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 26996.
WC200171042-A2.
                                                                                                                                                 ADI26064 standard; cDNA; 3430 BP.

Human cDNA encoding protein that promotes STAT6 activation #15
WC2003104277-A2.
                                                                                             10.0%; Score 27.4; DB 10; Length 3427; 55.9%; Pred. No. 6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.0%; Score 27.4; DB 12; Length 3617; 50.4%; Pred. No. 6.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 10.0%; Score 27.4; DB 4; Length 3615;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity 45.0%; Pred. No. 6.1e+02; Length 3615;
                                                                                                                                                                                                                                              Length 3430,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 4; Length 3614; 45.0%; Pred. No. 6.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                          10.0%; Score 27.4; DB 4; Length 3583; 52.1%; Pred. No. 6.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 3615;
                   Human STAT6-activating protein-encoding cDNA, SEQ ID NO:73.0200296943-A1.05-0202. (ASAH ) ASAHI KASEI KOGYO KK.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAI99443 standard; DNA; 3615 BP.
Human excretory related polynucleotide SEQ ID NO 1207.
WC200155313-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAI64088 standard; cDNA; 3615 BP.
Human bladder related polynucleotide, SEQ ID NO: 121
WC200159064-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 1369
ID ADF71686 standard; DNA; 3615 BP.
DE Human bladder associated antigen #16 genomic DNA #2.
PN US2003199008-A1.
                                                                                                                                                                                                                                              Score 27.4; DB 12;
Pred. No. 6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 MOLOULE
16-AUG-2001.
(HUMAL) HUMAN GENOME SCI INC.
HERY MATCh
10.0%; Score 27.4; DB 4;
ery Match
10.0%; Pred. No. 6.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human cDNA sequence SEQ ID NO:17248.
EP1074617-A2.
   ADG10483 standard; cDNA; 3427 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                 AAH17677 standard; cDNA; 3614 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADG31189 standard; DNA; 3617 BP.
Novel mouse gene #14.
WO2003089644-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ADG31187 standard; DNA; 3646 BP.
                                                                                                                                                                                      WO2003103.
18-DEC-2003.
(ASAH ) ASAHI KASEI KK.
- Match 10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      23-OCT-2003.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                (RIKE ) RIKEN KK.
(DNAF-) DNAFORM KK.
(MITU ) MITSUBISHI CHEM CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          30-OCT-2003.
(RIKE) RIKEN KK.
(DNAF-) DNAFORM KK.
(MITU ) MITSUBISHI CHEM CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       07-FEB-2001.
(HELI-) HELIX RES INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      .2-A2.
.2001.
.ry Match
Best Local Similarity :RESULT 1366
ID AAH17677 stand**
DE Human cDNA
PN EP10746*
PP 07.**
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Lot.

Lot.

Lot.

Lot.

Lot.

Lot.

Lot.

Best Local Similarity RESULT 1367

ID AA199443 stand*

DE Human excr*

PN WC2001**

PD 02.*

PA
                                                                                                                                                                                                                                           Query Match
Best Local Similarity
RESULT 1362
ID ABL10838 standard; cl
DE Drosophila melanogast
PN WC20171042-A2.
PD 27-SEP-2001.
PA (PEKE) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
                                                                                                          Best Local Similarity RESULT 1364
                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
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Length 4472;

Length 4507;

Length 4507;

#3.

Length 4507;

Length 5516;

50.4%; Pred. No. 7.9e+02;

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Best Local Similarity
RESULT 1390
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   30-JUL-1997
                                                                                                                                            Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               RESULT 1397
                                                                                                                                                                                                                                                                                                                                                                                                                                                        DE DE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAZ45239 standard; DNA; 6463 BP.
DNA encoding angiogenesis-associated protein which binds plasminogen.
W09966038-Al.
23-DEC-1999.
CPHAA ) PHARMACIA & UPJOHN AB.
10.0%; Score 27.4; DB 3; Length 6463;
Local Similarity 62.3%; Pred. No. 7.5e+02;
                                                                                                                                                                                                                                                                                                                                    AAS45338 standard; DNA; 6025 BP.
Chemically pretreated genomic DNA associated with cell cycle #22
WO200168911-A2.
20-SER-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                sequence #306
                            10.0%; Score 27.4; DB 11; Length 5572; 62.3%; Pred. No. 7.2e+02;
                                                                                                                                                                                                                                                                                        Length 5952;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (EPIG-) EPIGENOMICS AG. 10.0%; Score 27.4; DB 6; Length 7434;
                                                                                                                                                           Length 5875;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 6161;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human immune system associated gene SEQ ID NO: 1812 WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAS46584 standard; DNA; 6167 BP.
Tumour suppressor gene derived chemically modified
WO200168912-A2.
SO-SEP-2001.
(EPIG-) EPIGENOMICS AG.
10.0%; Score 27.4; DB 4; Le
st Local Similarity 49.6%; Pred. No. 7.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human immune system associated gene SEQ ID NO: 595.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human immune system associated gene SEQ ID NO: 880.
WO200200928-A2.
                                                                                         Human immune system associated gene SEQ ID NO: 261.
WO200200928-A2.
                                                                                                                                                                                                                                                 (BPIG-) EPIGENOMICS AG. 10.0%; Score 27.4; DB 10; ery Match initarity 55.9%; Pred. No. 7.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AG.
10.0%; Score 27.4; DB 6;
49.6%; Pred. No. 7.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 27.4; DB 6;
Pred. No. 7.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                               ZOCULE SETGENOMICS AG.

10.0%; Score 27.4; DB 4;

t Local Similarity 48.4%; Pred. No. 7.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           MAZZUZZ

02-JAN-2002.

(BEG) EPIGENOMICS AG.

(ERY Match 10.0%; Score 27.4; DB 6;

ery Match 47.4%; Pred. No. 7.4e+02;
                                                                                                                                                         Score 27.4; DB 6;
Pred. No. 7.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAD28386 standard; DNA; 7434 BP.
Human chemically treated genomic DNA
WO200202809-A2.
                                                                                                                                                                                                     ADBS4002 standard; DNA; 5952 BP. PCNA genomic DNA region.
WO2003072821-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DNA; 6685 BP
                                                                           ВР
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABL32622 standard; DNA; 6161 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 BP
                                                                                                                                        AG.
10.0%;
65.6%;
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10.0%;
69.8%;
                                                                             DNA; 5875
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DNA; 6167
Query Match
Best Local Similarity
RESULT 1381
ID ARITALE
                                                                                                                           03-JAN-2002.
(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               03-JAN-2002.
(EPIG-) EPIGENOMICS
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(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 1383
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Best Local Similarity
RESULT 1385
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Best Local Similarity
RESULT 1389
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Best Local Similarity
RESULT 1388
                                                                                                                                                         Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 1384
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL33839 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABL32907 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
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AAL37254 standard, DNA, 17908 BP.
Human musculoskeletal system related polynucleotide SEQ ID NO 3619.
WO200155367-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       cDNA encoding novel human musculoskeletal system antigen #2586.
US2002147140-A1.
10-OCT-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 13038;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 15109;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 14254;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 17908
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 17908
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                                                                                                                                                                                                                                  Length 8043;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 8126;
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Human nervous system related polynucleotide SEQ ID NO
WO200159063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           AAL03779 standard; DNA; 17908 BP.
Human reproductive system related antigen DNA SEQ ID
WO200155320-A2.
JULY LO. AND TABLE AND TAB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  JT 1398
ABL31374 standard; DNA; 13038 BP.
Human immune system associated gene SEQ ID NO: 1247.
WO200200928-A2.
03-JAN-2002.
(EPIG.) EPIGENOMICS AG.
10.0%; Score 27.4; DB 6; Le
sery Match
10.0%; Pred. No. 9.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                    23-DEC-1999.

(VLAA-) VLAAMS INTERUNIVERSITAIR INST BIOTECHNOG.
ery Match

ort Local Similarity 59.7%; Pred. No. 8.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (HUMA-) HUMAN GENOME SCI INC.

10.0%; Score 27.4; DB 2;

t Local Similarity 53.2%; Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.0%; Score 27.4; DB 5; 62.3%; Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 27.4; DB 6;
Pred. No. 1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
Query Match
Best Local Similarity 50.4%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.0%; Score 27.4; DB 4; 50.4%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Staphylococcus aureus contig SEQ ID #44
BP786519-A2
                                                                                                                                                                                                                                                                                                                               AAZ29606 standard; DNA; 8126 BP.
Basic fragmentation vector, pDVO DNA.
WO9966059-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CDNA; 17908 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         DNA; 15698 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DNA; 15109 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAV74355 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABL34140 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best_Local Similarity
RESULT 1392
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
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Query Match
                                                                                                                                                                                              RESULT 1400
ID AAS12088 standard; DNA; 18636 BP.
DE Human tumour necrosis factor receptor-like genomic polynucleotide #1.
PN W0200155440-A1.
                                                                                 Human musculoskeletal system-associated genomic DNA - SEQ ID 3619
US2004009488-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 exons 1-4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 18636,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL70481 standard; DNA; 23683 BP.
Chemically treated cell signalling DNA sequence#186.
W0200208807-A2.
(BPJG-2002.
(BPJG-) EPIGENOMICS AG.
ery Match
10.0%; Score 27.4; DB 6; Length 23683;
st Local Similarity 57.6%; Pred. No. 1.2e+03;
                                                                                                                                                           Length 17908
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 6; Length 18636; 55.9%; Pred. No. 1.1e+03;
      10.0%; Score 27.4; DB 8; Length 17908; 50.4%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 22509;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 23683;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAS44506 standard; DNA; 31529 BP.
Human LEKTI DNA clone CIT978SKB_94F21 contig 8, SPINK5 exon 5.
                                                                                                                                                                                                                                                                                                                   10.0%; Score 27.4; DB 5; Length 18636; 55.9%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 24183;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAS44505 standard; DNA; 22509 BP.
Human LEKTI DNA clone CIT978SKB_94F21 contig 11, SPINKS
WO200164747-A1.
                                                                                                                                                                                                                                                                                                                                                                                                #
                                                                                                                                                                                                                                                                                                                                                                        ABQ79004 standard; DNA; 18636 BP.
Human tumour necrosis factor gene from clone HEOQR40
US2002086820-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     USCULO.

USCULO.

(HUMA-) HUMAN GENOME SCI INC.

ery Match

10.0%; Score 27.4; DB 10;

ery Match

55.9%; Pred. No. 1.1e+03;
                                                                                                                       15-JAN-2004.
A (HUMA-) HUMAN GENOME SCI INC.
Ouery Match
Best Local Similarity 50.4%; Pred. No. 1.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           O'-SEP-2001.
(ISIS-) ISIS INNOVATION LTD.
Query Match
10.0%; Score 27.4; DB 4;
Query Match
53.2%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Wolver ...

18-02-1 - 2001.

(EPG-) EPIGENOMICS AG.

10.0%; Score 27.4; DB 6;

tery Match

10.0%; Score 27.4; DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.0%; Score 27.4; DB 5; 55.9%; Pred. No. 1.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 1404

DE ABL34622 standard; DNA; 23683 BP.

EN WO20017736-A2.

PD 18-OCT_2001.

PD 18-OCT_2001.

PA (EPIG-) EPIGRNOMPTC AC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADC35307 standard; DNA; 18636 BP.
Human TNFR-like gene 2 genomic sequence #1.
US2003077703-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (UYAL-) UNIV ALLEGHENY HEALTH SCI. (UYJE-) UNIV JEFFERSON THOMAS. (UYOU-) UNIV OULU.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAS21771 standard; DNA; 24183 BP.
Human gene for collagen COL9A1.
US6265157-B1.
                                                            ADJ30992 standard; DNA; 17908 BP.
                                                                                                                                                                                                                                                                             02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                             RESULT 1402

ID ADC35307 stand**

PN US20030**

PN US20030**

PD 24**

RESULT 1402

ID ADC35307 stand**

PN US20030**

PN US20030**

PN PS
                                                                                                                                                                                                                                                                                                                                                                                                                               04-JUL-2002.
(ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 1401
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 1407
Query Match
Best Local Similarity
RESULT 1399
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
Best Local Similarity
RESULT 1403
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
RESULT 1405
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           24-JUL-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
                                                                                                                                                                                                                                                                                                                     Ouery Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          RESULT 1406
ID AAS21777
DE Human ge
PN US62651
PD 24-JUL-,
PA (UYJE-)
PA (UYJE-)
PA (UYJE-)
                                                                                                                                                                                                                                                                                                                                                                            ID
DE
PA
PA
PA
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ABS55699 standard; DNA; 54842 BP.
Bovine Claudin-16 deficiency associated polynucleotide seguence #2.
JP2002238570-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   RESULT 1411

ID AAS46787 standard; DNA; 61020 BP.

DB Tunour suppressor gene derived chemically modified sequence #513.

PD 20-SEP-2001.

PD 20-SEP-2001.

PA (EPIG-) EPIGENOMICS AG.

Query Match

Query Match

Chery Match

Cher
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           12-JUN-2003.
(AMHP ) WYETH HOLDINGS CORP.
10.0%; Score 27.4; DB 9; Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 100543;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADA13316 standard; DNA; 108359 BP.

Human fringe-like secreted protein gene, SEQ ID NO:3.

Query Match

D10.0%; Score 27.4; DB 9; Length 108359;

Best Local Similarity 48.4%; Pred. No. 2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 110000;
WO200164747-A1.

07-SEP-2001.

(ISIS-) ISIS INNOVATION LTD.

(ISIS-) ISIS INNOVATION LTD.

10.0%; Score 27.4; DB 4; Length 31529;

(ery Match and S3.2%; Pred. No. 1.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAF28535 standard; DNA; 45613 BP.
Genomic fragment #22.
W020079968-A2.
28-DEC-2000.
[INCY-] INCYTE GENOMICS INC.
ET MACh
St.58; Pred. No. 1.5e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 37973;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 82938;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 65589;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 54842;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADB12064 standard; DNA; 1754382 BP.
Alloiococcus otitis entire genome sequence SEQ ID NO:6651.
WO2003048304-A2.
                                                                                                                                                                                                                                                                                                            Human immune system associated gene SEQ ID NO: 2170 WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        27-AUG-2002.

(CHIK-) CHIKUSAN GIJUTSU KYOKAI SH.

(KACH-) KACHIKU KAIRYO JIGYODAN SH.

(DOKU-) DOKURITSU GYOSEI HOJIN KACHIKU KAIRYO CE.

(DOKU-) MATCH.

ST. ACC ST. 4; DB 6;

St. Local Similarity 59.7%; Pred. No. 1.6e+03;
                                                                                                                                                                                                                                                                                                                                                                                       03-JAN-2002.
(BPIG-) EPIGENOMICS AG.
ery Match 10.0%; Score 27.4; DB 6;
EL Local Similarity 55.9%; Pred. No. 1.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ID ABV72623 standard; DNA; 82938 BP.

DB Human transporter protein encoding gene.

Query Match

Best Local Similarity 75.6%; Pred. No. 1.9e+03;

RESULT 1414
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.4; DB 6; 52.1%; Pred. No. 2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 6; 53.2%; Pred. No. 1.7e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.0%; Score 27.4; DB 6; 50.4%; Pred. No. 2.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.0%; Score 27.4; DB 6; 53.2%; Pred. No. 2.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABS52816 standard; DNA; 100543 BP.
Genomic DNA encoding human secreted protein #4
WO200264626-A2.
                                                                                                                                                                                     Best Local Similarity RESULT 1408
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best_Local Similarity
RESULT 1413
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               (PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match
Best Local Similarity
RESULT 1417
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RESULT 1418
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      22-AUG-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Si
RESULT 1409
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                                                                                                                                                    Query Match
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Score 27.2; DB 10; Length 393;
Pred. No. 3.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (HUMA.) HUMAN GENOME SCI INC.
10.0%; Score 27.2; DB 12; Length 449;
it Local Similarity 53.8%; Pred. No. 3.3e+02;
                                                                                                                                                                                                      Length 359;
                                                       Length 317;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 414;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 449;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 449;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 451;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 451;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Colon adenocarcinoma related gene sequence SEQ ID NO:953 WO200194629-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             319
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           50
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human excretory related polynucleotide SEQ ID NO WO200155313-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAI64062 standard; cDNA; 449 BP.
Human bladder related polynucleotide, SEQ ID NO:
WO200159064-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      16-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
10.0%; Score 27.2; DB 4;
Lery Match 13.18; Pred. No. 3.3e+02;
                                                     Score 27.2; DB 8;
Pred. No. 2.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 27.2; DB 8;
Pred. No. 3.2e+02;
                                                                                                                                                                                  (USSH ) US DEPT HEALTH & HUMAN SERVICE.
ry Match 10.0%; Score 27.2; DB 2;
t Local Similarity 50.0%; Pred. No. 3.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 27.2; DB 6;
Pred. No. 3.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 27.2; DB 6;
Pred. No. 3.3e+02;
                                                                                                                                                                                                                                          Human polynucleotide sequence SEQ ID NO:816.
MO2003080795-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABN67263 standard; DNA; 489 BP.
Streptococcus polynucleotide SEQ ID NO 2439.
WO200234771-A2.
                                                                                                                       Human brain Expressed Sequence Tag EST01293 W09316178-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 1435
ID ADF71615 standard; CDNA; 449 BP.
DE Human bladder associated antigen cDNA #40.
PN US2003199008-A1.
PD 23-CCT-2003.
PA (HUMA-) HUMAN CENAMA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Gene #1387 used to diagnose liver cancer.
                                                                                                                                                                                                                                                                                                                                                                                                                           Prokaryotic essential gene #5587
WO200277183-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          CDNA; 449 BP
                                                                                                             DNA; 359 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ABL62616 standard; DNA; 451 BP.
                                                                                                                                                                                                                                                                                                                                                                                                          DNA; 414 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                  WOZUCZ...
03-007-2002.
(ELIT-) ELITRA PHARM INC.
10.0%;
                                                     10.0%;
50.8%;
                                                                                                                                                                                                                                                                                                                                                    10.0%;
51.7%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PD 11-APR-2002.

PA (GENE-) GENE LOGIC INC.

Bet I Local Similarity 48.7%;

RESULT 1438
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         13-DEC-2001.
(AVAL-) AVALON PHARM.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 1436
                                                                                                                                                                                                                 Best Local Similarity
RESULT 1431
                                                                   Best Local Similarity
RESULT 1430
                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 1433
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
                                                                                                             AAQ61275 standard;
                                                                                                                                                                                                                                                                                                           02-OCT-2003.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAI98875 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                            ACA23930 standard;
                                                                                                                                                                   19-AUG-1993
                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
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                                                       Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABX42543 standard; cDNA; 317 BP.
Bovine EST associated with lactation/muscle/fat deposition #7708.
US2002137139-Al.
26-SEP-2002.
                                      Length 110000;
                                                                                            Length 110000,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 181343;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 192427;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 200620;
                                                                                                                                                  Length 110000
                                                                                                                                                                                                AAF25833 standard; DNA; 130480 BP.
R. marinus bacteriophage RM387 genomic DNA.
W020075335-A2.
14-DEC-2000.
(DECO-) DECODE GENETICS EHF.
ery Match
St Local Similarity 49.68; Pred. No. 2.2e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              27-JUN-zvuz.
(GENE JOGIE LOGIC INC.
(PROC. ) PROCTER & GAMBLE CO.
10.0%; Score 27.4; DB 6; Length 160771;
rry Match 10.0%; Pred. No. 2.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 147724;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 174424;
                                                                                                                                                                                                                                                                                                                                                  ABKR81566 standard; cDNA; 147724 BP.
Human cDNA differentially expressed in granulocytic cells #137
WO200228999-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human presynaptic cytomatrix protein, REPS2, genomic sequence WO2004044164-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       98
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABQ88179 standard; cDNA; 160771 BP.
Human osteoblast differentiation related cDNA SEQ ID NO
WO200250301-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADQ19573 standard; DNA; 181343 BP.
Human soft tissue sarcoma-upregulated DNA - SEQ ID 2392.
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADL13825 standard; DNA; 192427 BP.
Ostecarthritis-associated polymorphic nucleotide #357.
WO2003054166-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Ovary cancer related gene sequence SEQ ID NO:6459 WO200194629-A2.
13-DEC-2001.
(AVAL-) AVALON PHARM.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (PROT-) PROTEIN DESIGN LABS INC.
(PROT-) PROTEIN DESIGN LABS INC.
10.0%; Score 27.4; DB 12;
(ery Match
10.0%; Pred. No. 2.4e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                03-UUL-2003.
(INCY-) INCYTE GENOMICS INC.
(INCY-) INCYTE GENOMICS INC.
10.0%; SCOIE 27.4; DB 10;
GETY MAICh 12.5e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.0%; Score 27.4; DB 12; 57.6%; Pred. No. 2.5e+03;
                                    Score 27.4; DB 12;
Pred. No. 2.1e+03;
                                                                                        Score 27.4; DB 12;
Pred. No. 2.1e+03;
                                                                                                                                                Score 27.4; DB 12;
Pred. No. 2.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                   GENE LOGIC INC.

th. 10.0%; Score 27.4; DB 6;

Similarity 59.7%; Pred. No. 2.3e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.4; DB 6; 53.2%; Pred. No. 2.4e+03;
   Pred. No. 2.1e+03;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL68122 standard; DNA; 174424 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADO56277 standard; DNA; 200620 BP
                                                                                                                                              10.0%;
50.4%;
                                  10.0%;
50.4%;
                                                                                        10.0%;
50.4%;
 53.2%;
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(SEQU-) SEQUENOM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 1425
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 1423
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Best Local Similarity
RESULT 1424
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RESULT 1428
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RESULT 1427
                                                                       RESULT 1420
Query Match
Best Local Similarity
                                                                                                                                              Query Match
Best Local Similarity
RESULT 1422
ID AAF25833 standard;
   Local Similarity
                                    Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (BYAT/) BYATT J C.
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11-APR-2002

27-JUN-2002

Query Match

Query Match

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(SMIK ) SMITHKLINE BEECHAM CORP. (SMIK ) SMITHKLINE BEECHAM PLC. Query Match 10.0%; Scc
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(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                    10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.0%;
55.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     10.0%;
58.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
RESULT 1452
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 1454
                                                                                                                                                                                               Best Local Similarity RESULT 1449
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
RESULT 1453
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 RESULT 1451
ID AAI64132 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
Best Local Similarity
RESULT 1455
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 1456
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (RIKE ) RIKEN KK.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              EP1033405-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         24-JUN-1999
   07-MAR-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       04-JUN-1998
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 30-MAY-2003
                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADN13758 standard; cDNA; 582 BP.
Human prostate/colon/lung/breast cancer-related cDNA 1273, SEQ:1273.
WO2004039943-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABQ31082 standard; DNA; 784 BP.
Oligonuclectide for detecting cytosine methylation SEQ ID NO 17673.
WO200218632-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABN91349 standard; DNA; 699 BP.
Staphylococcus epidermidis ORF nucleic acid sequence SEQ ID NO:812.
US6380370-B1.
                                                                                                          ADNIZ790 standard; cDNA; 517 BP.
Human prostate/colon/lung/breast cancer-related cDNA 305, SEQ:305.
WC2004039943-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human prostate/colon/lung/breast cancer-related cDNA 248, SEQ:248.
WQ2004039993-A2.
                                                                                                                                                                                                     10.0%; Score 27.2; DB 12; Length 517; 50.4%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.0%; Score 27.2; DB 12; Length 582; 50.4%; Pred. No. 3.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 582;
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                                                                                                                                                                                                                                                                                                                                                  Length 520;
                                                         Length 489;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 530;
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PA (GENO-) GENOME THERAPEUTICS CORP.
Query Match
Best Local Similarity 52.7%; Pred. No. 3.9e+02;
ID 80-3031082 standard; DNA; 784 BP
DE Oligonucleotide for 3.94 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.0%; Score 27.2; DB 12; 50.4%; Pred. No. 3.7e+02;
                                                         10.0%; Score 27.2; DB 6; 53.8%; Pred. No. 3.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.2; DB 6; 64.1%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ry Match 10.0%; Score 27.2; DB 5;
t Local Similarity 53.9%; Pred. No. 3.9e+02;
                                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.2; DB 3; 54.1%; Pred. No. 3.5e+02;
                                                                                                                                                                                                                                                                                                            Best Local Similarity 10.0%; Score 27.2; DB 3 RESULT 1441

ID ABW87968 standard; CDNA; 530 BP.
DE Human colon concer related CDNA SEQ ID NO 1279.
PD 01.AUG-2002.
PA (CORI-) COLO.
                                                                                                                                                                                                                                                       AAA26713 standard; DNA; S20 BP.
Candida albicans polynucleotide sequence #51.
BP982401-A2.
(JAAR-2000.
(JANC ) JANSSEN PHARM NV.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABV21872 standard; cDNA; 683 BP.
Human prostate expression marker cDNA 21863
WC200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Human prostate expression marker cDNA 27693 WO200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ) 23-AUG-2001.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

Query Match 10.0%; Score 27.2; D
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABV27702 standard; cDNA; 683 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADN12733 standard; cDNA; 582 BP
02-MAY-2002.
(CHIR-) CHIRON SPA.
(GENO-) INST GENOMIC RES.
                                                                                                                                      ...R ) CHIRON CORP.
..ery Match
Best Local Similarity 5
RESULT 1440
ID AAA26713 stand**
DE Candida al**
PD 01-"
PA 01-"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            "-43-A2.
"AR ) CHIRON CORP
"CET MATCH
Best Local Similarity L
RESULT 1443
ID ADN13758 stand**
DE Human prost*
PN W02004^*
PD 13-**
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity (RESULT 1442 ID ADM12733 standard; CDP DE Human prostate/colon/1PN WO2004019943-A2.
PD 13-MAY-2004.
PA (CHIR ) CHIRON CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        JO-A2.

LLL-) MILLENNIUM

LLL-) MILLENNIUM

LAY MACCA Similarity

RESULT 1445

ID ABV27702 stand**

DB Human prost*

PN W02001f*

PD 23-*

PA
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(CHIR ) CHIRON CORP.
                                                                      Best Local Similarity
RESULT 1439
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                                                          Query Match
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ABO31083 standard; DNA; 784 BP.
Oligonucleoride for detecting cytosine methylation SEQ ID NO 17674.
WO200218632-A2.
(RPIG-) EPIGENOMICS AG.
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Extended human secreted protein coding sequence, SEQ ID NO. 74.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 957;
                         Length 784;
                                                                                                                                                                                                                       Length 784;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 A199487 standard; DNA; 957 BP.

Human excretory related polynucleotide SEQ ID NO 1251
W0200155313-A2.
02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
st Local Similarity 53.8%; Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AA164132 standard; CDNA; 957 BP.
Human bladder related polynucleotide, SEQ ID NO: 165
WO200159064-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         #
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Streptococcus pneumoniae polypeptide coding region.
WO9823631-A1.
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Arabidopsis thaliana DNA fragment SEQ ID NO: 1510.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%; Score 27.2; DB 12; 53.8%; Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADF71730 standard; DNA; 957 BP.
Human bladder associated antigen #40 genomic DNA
US2003199008-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADB23199 standard; DNA; 1000 BP.
Environmental stress-responsive promoter, SEQ ID
WO2003044190-Al.
PA (EPIG-) EPIGENOMICS AG.
Query Match
10.0%; Score 27.2; DB 6;
Best Local Similarity 58.8%; Pred. No. 4.1e+02;
RESULT 1448
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16-AUG-2001.
(HTMA.) HUMAN GENOME SCI INC.
(HTMA.) HUMAN GENOME SCI INC.
10.0%; Score 27.2; DB 4;
ery Match
' c'milarity 53.8%; Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           10.0%; Score 27.2; DB 2; 50.8%; Pred. No. 4.4e+02;
                                                                                                                                                                                                                                                                                                                                                             PD 07-FEB-2001.

PA (HELL-) HELIX RES INST.

QUENTY MATCh 10.0%; SCORE 27.2; DB 4;
BEST LOCAL Similarity 55.2%; Pred. No. 4.1e+02;
RESULT 1450
                                                                                                                                                                                                                       Score 27.2; DB 6;
Pred. No. 4.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 27.2; DB 9;
Pred. No. 4.4e+02;
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Pred. No. 4.5e+02;
                                                                                                                                                                                                                                                                                 AAH03690 standard; cDNA; 788 BP.
Human cDNA clone (5'-primer) SEQ ID NO:525.
EP1074617-A2.
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14-NOV-2002.
(HUMA-) HUMAN GENOME SCI INC.
10.0%; Score 27.2; DB 10; Length 2520;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 2000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 1994;
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Length 1747;
                                                                                                                                                                                                                                                                                                            Length 1854;
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                                                                                                                                                    Length 1785
                                                                                                                                                                                                             ADA21174 standard; cDNA; 1854 BP.
Human secreted protein SECP-28 encoding cDNA SEQ ID NO:79-
WO20031668943-A2.
                                                    ABL17101 standard, DNA, 1785 BP.
Drosophila melanogaster genomic polynucleotide SEQ ID NO
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Disease treating protein complex-derived gene #1226
EP1338608-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABQ80054 standard; cDNA; 2399 BP.
Corn granule-bound starch synthase coding sequence CA2384605-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 27.2; DB 10;
Pred. No. 5.7e+02;
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03-0002.

(DUPO ) DU PONT DE NEMOURS & CO E I.

ery Match 10.0%; Score 27.2; DB 8;

corr 27.2; DB 8;

corr 27.2; DB 8;
Score 27.2; DB 2;
Pred. No. 5.4e+02;
                                                                                                                                                                                                                                                                     PD 21-AUG-2003.

PA (INCY-) INCYTE GENOMICS INC.

Query Match

10.0%; Score 27.2; DB 9;

Best Local Similarity 58.8%; Pred. No. 5.6e+02;

RESULT 1468
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.0%; Score 27.2; DB 3; 50.0%; Pred. No. 6.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.0%; Score 27.2; DB 2; 51.7%; Pred. No. 5.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         10.0%; Score 27.2; DB 8; 50.0%; Pred. No. 6.2e+02;
                                                                                                                                                    Score 27.2; DB 4;
Pred. No. 5.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAA76420 standard; cDNA; 2520 BP.
Human secreted protein gene 40 SEQ ID NO:50.
WO200035937-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADD37651 standard; cDNA; 2520 BP.
Human secreted protein encoding sequence
WO200290526-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human secreted protein encoding cDNA. W02002102993-A2.
                                                                                                                                                                                                                                                                                                                                                                 AAT09085 standard; cDNA; 1994 BP. Acharina fulica cDNA. DP07233195-A. CS-SEP-1995. (SUNR ) SUNTORY LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ACC61635 standard; DNA; 2000 BP. Gene sequence #SEQ ID 2052.
EP1258494-A1.
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10.0%;
                                                                                                                                                    10.0%;
61.1%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          10.0%;
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53.8%;
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(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      20-NOV-2002.
(CELL-) CELLZOME AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                27-AUG-2003.
(CELL-) CELLZOME AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 1470
                                                                                                               27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 1469
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity
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Best Local Similarity
RESULT 1474
Query Match
Best Local Similarity
RESULT 1466
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Local Similarity
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                                                                                                                                                                        Local Similarity
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CDNA encoding human alpha 2, 3-sialyltransferase (ST3 GalVI) 9-13.
CN1329158-A.
                                                                                                                                                                        10.0%; Score 27.2; DB 12; Length 1153; 47.6%; Pred. No. 4.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 1428;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        10.0%; Score 27.2; DB 8; Length 1359; 58.8%; Pred. No. 5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 1467;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   PD 02-JAN-2002.

PA (SHAN-) SHANGHAI BIODOOR GENE DBV CO LTD.

Query Match 10.0%; Score 27.2; DB 6; Length 1633;

Best Local Similarity 53.8%; Pred. No. 5.3e+02;

RESULT 1464
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10.0%; Score 27.2; DB 3; Length 1723; 55.2%; Pred. No. 5.4e+02;
                      Length 1153;
                                                                                                                                                                                                                                                                                                                                Length 1275;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 1467;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABZ14297 standard; DNA; 1467 BP.
Arbidopsis thaliana stress regulated gene SEQ ID NO 2102.
WO200216555-A2.
28-FBB-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAC51519 standard; DNA; 1723 BP.
Arabidopsis thaliana DNA fragment SEQ ID NO: 68821.
EP1033405-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  12-AUG-2003.
(GENO-) GENOME THERAPEUTICS CORP.
ery Match
ery ** Score 27.2; DB 10;
ery ** Score 27.2; DB 10;
ery ** Score 27.2; DB 10;
                                                                                                                                                                                                                                                                                                                                10.0%; Score 27.2; DB 8; 47.6%; Pred. No. 4.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             10.0%; Score 27.2; DB 6; 55.2%; Pred. No. 5.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     03-JAN-2003.

(SYGN ) SYNGENTA PARTICIPATIONS AG.

(SYGN ) SYNGENTA PARTICIPATIONS AG.

ery Match

Ery Match 10.0%; Score 27.2; DB 8;

ery Match 55.2%; Pred. No. 5.1e+02;
                    10.0%; Score 27.2; DB 2; 47.6%; Pred. No. 4.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Arabidopsis thaliana gene, SEQ ID 484.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (SCRI ) SCRIPPS RES INST.
(SYGN ) SYNGENTA PARTICIPATIONS AG.
                                                                           ADP18877 standard; cDNA; 1153 BP. Human secreted polynucleotide #133.0204110939-A1. (GEST ) GENSET SA.
                                                                                                                                                                                                                                                                                                                                                                           Prokaryotic essential gene #14115.
WO200277183-A2.
                                                                                                                                                                                                                           17 1458

Prokaryotic essential gene #10676.

020027183-A2.

03-0C1-2002.

(ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADF02388 standard; DNA; 1428 BP.
Bacterial polynucleotide #2673.
US6605709-B1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DNA; 1747 BP. (rib-1).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADA68572 standard; DNA; 1467 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (ELIT-) ELITRA PHARM INC.
                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 1459
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
RESULT 1461
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity RESULT 1462
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Best Local Similarity
RESULT 1463
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 1465
                                                                                                                                                                                        Best Local Similarity RESULT 1458
                    Query Match
Best Local Similarity
RESULT 1457
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAQ62810 standard;
Riboflavin 1 gene
DE4238904-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 26-MAY-1994.
(BADI ) BASF AG.
  (GEST ) GENSET.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           06-SEP-2000.
                                                                                                                                                                              Query Match
                                                                                                                                                                                                                                                                                                                                  Query Match
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ID DE PN

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27-DEC-2002.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
lery Match
lery Match
10.0%; Score 27.2; DB 10; Length 2520;
                                                                                                                                                                                                                                                                                                                                                S. epidermidis genomic polynucleotide sequence SEQ ID NO:4062. WC200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  RESULT 1483
ID AAH54424 standard; DNA; 3673 BP.
DE S. epidermidis genomic polynucleotide sequence SEQ ID NO:3788.
PN WO200134809-A2.
                                                                                                                                                                                                                                                                             Length 2761;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAH54829 standard; DNA; 3041 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:4193
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAHS4143 standard; DNA; 3349 BP.
S. epidermidis genomic polynucleotide sequence SEQ ID NO:3507
WO200134809-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     | 17-MAY-2001.
| (GLAX) | GLAXO GROUP LTD.
| 10.0%; Score 27.2; DB 4; Length 3673;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 3048;
                                                                                                                                                                                                                                                                                                                                                                                                                         Score 27.2; DB 4; Length 2982;
Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 3041;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.0%; Score 27.2; DB 6; Length 3096; 50.0%; Pred. No. 6.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 3349,
                                                                                                                                                                              AAC85567 standard; cDNA; 2761 BP.
cDNA encoding CDIFF-27, Incyte ID No. 2573955CB1
WC200119860-A2.
                                                                                                                                                                                                                                                      (INCY-) INCYTE GENOMICS INC.
ry Match
t Local Similarity 58.8%; Pred. No. 6.4e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    10.0%; Score 27.2; DB 4; 55.2%; Pred. No. 6.6e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 27.2; DB 6;
Pred. No. 6.7e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 27.2; DB 4;
Pred. No. 6.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 27.2; DB 4;
Pred. No. 6.6e+02;
50.0%; Pred. No. 6.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABA92165 standard; DNA; 3057 BP.
Arabidopsis ATP sulfurase gene (APS1) promoter.
WO200216621-A2,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABN68454 standard, DNA, 3096 BP.
Streptococcus polynucleotide SEQ ID NO 4821.
WO200234771-A2.
                                  ADA56221 standard; DNA; 2520 BP.
Gene encoding human secreted protein #400.
WC2002102994-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAH18354 standard; cDNA; 3048 BP.
Human cDNA sequence SEQ ID NO:18387.
EP1074617-A2.
                                                                                                                                                                                                                                                                                                                           AAH54698 standard; DNA; 2982 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%;
51.7%;
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50.0%;
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A (CHIR-) CHIRON SPA.
A (GENO-) INST GENOMIC RES
Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
                                                                                                                                                                                                                                                                                                                                                                                   17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           17-MAY-2001.
(GLAX ) GLAXO GROUP LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               07-FEB-2001.
(HELI-) HELIX RES INST.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
RESULT 1480
                                                                                                                            Query Match
Best Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity RESULT 1482
Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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(RHOB-) RHOBIO.
                                                                                                                                                                                                                                       22-MAR-2001
                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
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                                                                                                                                                          RESULT 1476
1D AAC85567
DE CDNA enc
PN WO200119
PD 22-MAR-2
PA (INCY-)
                                                                                                                                                                                                                                                                                                                                                                                                                                                             RESULT 1478
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               RESULT 1475
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ABT23200 standard; DNA; 4384 BP.
Human protein modification + maintenance molecule DNA SEQ ID No 29.
W02003000844-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABLO8292 standard; cDNA; 4742 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 19358.
WO200171042-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Drosophila metanogater expressed polynucleotide SEQ ID NO 1046. PERS PE CORP NY.
                                                                                        10-2004.
(PROT-) PROTEIN DESIGN LABS INC.
(ery Match 10.0%; Score 27.2; DB 12; Length 3729;
                                                                                                                                                                                                                                                                                   Length 3905;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 3905;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   10.0%; Score 27.2; DB 4; Length 4936; 55.2%; Pred. No. 7.9e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 4920;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 4384
                                                           Human soft tissue sarcoma-upregulated DNA - SEQ ID 7216.
WO2004048938-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             PA (FDA...)

Query Match

Best Local Similarity 53.8%; ...

RESULT 1490

ID ABL34284 standard; DNA; 4920 BP.

DE Human inmune system associated gene SEQ ID NO: 2257.

DO 03-JAN-2002.

PD 03-JAN-2002.

PA (BPIG-) EPIGENOMICS AG.

"...orv Match 10.0%; Score 27.2; DB 6; Leng'

"...orv Match"...ilarity 52.7%; Pred. No. 7.9e+02;
                                                                                                                                                                                       AAX13036 standard, DNA, 3905 BP.
Enterococcus faecalis genome contig SEQ ID NO:99.
WO9850555-A2.
                                                                                                                                                                                                                                       12-NOV-1998.

(HUMA-) HUMAN GENOME SCI INC.

(HTMA-) HUMAN GENOME SCI INC.

10.0%; Score 27.2; DB 2;

ery Match 10.0%; S9.5%; Pred. No. 7.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 27.2; DB 4;
Pred. No. 7.5e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 27.2; DB 6;
Pred. No. 7.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      .2; DB 8;
. 7.6e+02;
       52.7%; Pred. No. 7.1e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAS44657 standard; DNA; 4936 BP.
Mouse full-length polynucleotide sequence #82.
WO200164834-A2.
                                                                                                                                                                                                                                                                                                                                   ABS98831 standard, DNA, 3905 BP.
Enterococcus faecalis contig sequence #99.
US2003120116-A1.
29-AUG-2002.
(KUNS), KUNSCH C A.
(DILL/) DILLON P J.
(BARA/) BARASH S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 27.2
Pred. No.
                                      ADQ24396 standard; DNA; 3729 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         BP.
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Genomic DNA of the CCBP2 gene.
W0200232926-A2.
25-APR-2002.
(GENA-) GENAISSANCE PHARM INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       10.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       MOSTAL 2003.
(INCY-) INCYTE GENOMICS INC.
lery Match
10.0%;
Best Local Similarity RESULT 1484
                                                                                                                                                Best Local Similarity RESULT 1485
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
RESULT 1487
ID ABL02188 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity RESULT 1488
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
RESULT 1489
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Ouery Match
Best Local Similarity
RESULT 1492
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           07-SEP-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                   Query Match
                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
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ABLO7648 standard; cDNA; 5970 BP.
Drosophila melanogaster expressed polynucleotide SEQ ID NO 17426.
27-SEP-2001.
(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                            AAS46462 standard; DNA; 5686 BP.
Tumour suppressor gene derived chemically modified sequence #184
W0200168912-A2.
20-SEE-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity 64.1%; Pred. No. 8.38+02;
RESULT 1496
ID ABL19820 standard; DNA; 5690 BP.
DE Drosophila melanogaster genomic polynucleotide SEQ ID NO 10933.
PN WO200171042-A2.
                                                                                                                                                                            10.0%; Score 27.2; DB 10; Length 5382; 56.8%; Pred. No. 8.2e+02;
                                                                                                                                                                                                                                                                                                                                                         Length 5468;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    LT 1493

AR31432 standard; DNA; 6099 BP.

Signal transduction associated gene modified DNA #138.

W0200026-A2.

W0200026-A2.

(EPTAN-2002.

(EPTG-) EPIGENOMICS AG.

ETY Match

St. Local Similarity 53.8%; Pred. No. 8.5e+02;
  10.0%; Score 27.2; DB 6; Length 5090; 67.9%; Pred. No. 8e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 5970;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      10.0%; Score 27.2; DB 4; Length 5690; 64.1%; Pred. No. 8.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA; 6099 BP.
cell signalling DNA sequence#144.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL12878 standard; DNA; 5994 BP.
Human immune system associated gene SEQ ID NO: 851.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                       10.0%; Score 27.2; DB 10; 56.8%; Pred. No. 8.2e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               20.5br-zour.
(BPIG-) BPIGENOMICS AG.
ry Match 10.0%; Score 27.2; DB 4;
r Tocal Similarity 64.1%; Pred. No. 8.3e+02;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       MCACCLOS.
(PDIG-) EDIGENOMICS AG.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               10.0%; Score 27.2; DB 4; 53.8%; Pred. No. 8.5e+02;
                                            Est Local Similarity 56.8%; Pred. N RESULT 1494
ID ADEG3061 standard; DNA; 5468 BP.
DE Human gene M80340, SEQ ID NO 8996. PN W02003016475-A2. PD 27-FEB-2003. PA (GEHO ) GEN HOSPITAL CORP.
PA (FARE ) BAYER AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AG.
10.0%;
53.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 1500
ID ABL70397 standard; DNI
DE Chemically treated ce-
PN WO200202807-A2.
PD 10-JAN-2002.
PA (EPIG-) EPIGENOMICS AC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10-JAN-2002
(EPIG-) EPIGENOMICS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
RESULT 1498
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Best Local Similarity
Query Match
Best Local Similarity
RESULT 1493
                                                                                                                                                                                                                                                                                                                                                       Query Match
Best Local Similarity
RESULT 1495
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(PEKE ) PE CORP NY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity
RESULT 1497
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Best Local Similarity
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AC129793 Rattus no AC112376 Rattus no AC097216 Rattus no AF172282 Oryza sat AR355053 Sequence AL401438 T7 end of AF194946 Escherich AY25001 Escherich X54971 Xiphophorus U35049 Arabidopsis AX281516 Sequence	AX34540 sequence AL110495 Caenorhab AC116100 Dictyoste AC06819 Homo saphi AC014345 Drosophil AP004644 Oryza sat AC118284 Oryza sat	AC102011 Mus muscu AC102011 Mus muscu AC114905 Mus muscu AC008928 Home sapi AL356240 Human DNA AL646048 Mouse DNA AC11167 Mus muscu	AC13435 House Sapi AC134433 Mus muscu AC124960 Medicago AL60655 Oryza sat AL60665 Oryza sat AL60665 Oryza sat AC04811 Sus scrof AC16841 Mus muscu AC14817 Macaca mu	AC119168 Mus muscu AL035697 Human DNA BX510372 Zebrafish AP005687 Oryza sat AC010835 Homo sapi AP00467 Oryza sat AC105391 Homo sapi AP001883 Homo sapi AC024859 Caenorhab CR388171 Danio rer AC110822 Mus muscu	AP005BT6 Curyas ast AC090051 Homo sapi AC118826 Rattus no AC192711 Kenopus t AC09188 Homo sapi AC19238 Mus muscu AC115598 DictyOste AC020605 Homo sapi AC036150 Homo sapi AC036150 Homo sapi AC036150 Sus scrof AC002451 Homo sapi	AC125130 Mus muscu AC12028 Rattus no BX855617 Zebratish AL672023 Mouse DNA AC04957 Homo sapi AC107021 Homo sapi AC107021 Homo sapi AC107021 Homo sapi AC12304 Homo sapi AC123035 Mus muscu AC123035 Mus muscu AC192673 Homo sapi AC12035 Mus muscu AC12530 Zebratish AC126845 Homo sapi
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31.4 11.5 150634 2 ACI13371 ACC04289 AC	31.4 11.5 216208 3 31.4 11.5 217446 2 31.4 11.5 220618 2 31.4 11.5 220618 2 31.4 11.5 220618 2 31.4 11.5 220618 2 31.4 11.5 231759 2 31.4 11.5 231759 2 31.4 11.5 234304 5 31.4 11.5 234304 5 31.4 11.5 234304 5 31.4 11.5 234304 2 31.4 11.5 234304 2 31.4 11.5 234304 2 31.4 11.5 243642 2 31.4 11.5 243642 2 31.4 11.5 244385 2 31.4 11.5 246310 2 31.4 11.5 246310 2 31.4 11.5 256643 2 31.4 11.5 256643 2 31.4 11.5 256643 2 31.4 11.5 256643 2 31.4 11.5 256643 2 31.4 11.5 256643 2 31.4 11.5 256643 2 31.4 11.5 256643 3 31.4 11.5 256643 3 31.4 11.5 256643 3 31.4 11.5 256643 3 31.4 11.5 256643 3 31.4 11.5 256643 3 31.4 11.5 256643 3 31.4 11.5 256644 3 31.4 11.5 256644 3 31.4 11.5 256644 3 31.4 11.5 26684 3 31.4 11.5 26684 3 31.4 11.5 277968 2 31.4 11.5 293218 2 31.4 11.5 293218 2 31.4 11.5 293218 2 31.4 11.5 293218 2	31.2 11.9 31031.3 31.2 31.2 31.2 31.2 31.2 31.2 31.2
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31.2 11.4 181841 10 AC138792 AC128001 Rattus muscu ac12801 2 AC128001 Rattus mo ac128001 Rattus mo ac128001 Rattus mo ac12800 2 AC128001 Rattus mo ac12801 2 AC128001 Rattus mo ac12801 2 AC12802 2 AC13468 Mus muscu ac12812 11.4 183279 2 AC13468 Mus muscu accordance	AR252626 AR252626 AR252626.1 GI:27300534 Unknown. Unclassified. 1 (bases 1 to 713) Winterbottom, J.M., Shimp, L., Boyce, T.M. and Kaes, D. Implant, method of making same and use of the implant for the treatment of bone defects Patent: US 6478825-A 376 12-NOV-2002; Location/Qualifiers 1. 713 /organism="unknown" /mol_type="genomic DNA"	y Match Local Similarity 100.0%; Score 273; DB 6; Length 713; Local Similarity 100.0%; Pred. No. 2.66-65; hes 273; Conservative 0; Mismatches 0; Indels 0; Gaps 0; 1 ATGACATTTTTCTGTCACTATTATTGTTGTGAAGCTATTTGAAGAATCCAAT 60
114880 14480 144881 144881 144883 144886 14499 14496 14996 11496 11496 11496 11496 11496	RESULT 1 AR252626 LOCUS DEFINITION ACCESSION VERSION KEYWORDS SOURCE ORGANISM REFERENCE AUTHORS TITLE JOURNAL FEATURES SOURCE	Query Match Best Local Matches 27 Qy 61 Db 152 Qy 121 Db 212 Qy 181 Db 272 Qy 241 Db 332 Qy 332 RESULT 2 332
AP006139 Lotus cor AL356778 Human DNA AC079374 Rarbidops AC001645 Arabidops AL049660 Arabidops BX548058 Zebrafish AL590635 Human DNA BX927197 Danito rer AC084705 Arabidops AC108086 Homo sapi AC108086 Homo sapi AC004259 Human DNA s Continuation (4 of CX387919 Danito rer AC084519 Homo sapi AC004559 Human Chr AC06951 Homo sapi AC06951 Homo sapi AC06951 Homo sapi AC06951 Homo sapi AC11525 Homo sapi AC115278 MATUS SATUNGYIO AC149978 STRONGYIO AC149978 STRONGYIO	AC114594 Mus muscu AC09509 Homo sapi BX510325 Zebrafish AL121990 Human DNA AC026175 Homo sapi AC141740 Apis mell BX323067 Zebrafish BX323067 Zebrafish AC027196 Homo sapi AC07196 Homo sapi AC016301 Homo sapi AC01376 Homo sapi AC01376 Homo sapi AC01376 Homo sapi AC069532 Homo sapi AC069536 Rattus no AL908140 Mouse DNA AC10656 Homo sapi	AC023110 Home sapi AC023110 Home sapi AC023110 Home sapi AC120393 Mus muscu AC12744 Mus muscu AC12744 Mus muscu AC127546 Human DNA AC12693 Rattus no AC450346 Human DNA AC12693 Rattus no AC450346 Human Sapi BX314216 Danio rer AC011628 Mus muscu AC012164 Danio sapi BX314216 Danio sapi AC012693 Home sapi AC024479 Home sapi AC024479 Home sapi AC022012 Home sapi AC12773 Rattus no BX276188 Zebrafish AC114877 Home sapi
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Gaps

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Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
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Gao,W.Q., Gerritsen,M.E., Goddard,A., Godowski,P.J., Gurney,A.L.,
Sherwood,S., Smith,V., Stewart,T.A., Tumas,D., Watanabe,C.K.,
Wood,W.L. and Zhang,Z.
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                                                                                                                                                                                   1 ATGACATTTTTTCTGTCACTATTATTATTGTTGGTATGTGAAGCTATTTGGAGATCCAAT
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                                                                                                                                                                                                                                                                                                                                                CAACCCACACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACA
                                                                                                                                                                                                                                                                                                                                                                                    212 CAACCCACACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACA
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                                                                                                     Length 713;
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Pred. No. 2.6e-65;
; Mismatches 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 332 GCCTGGGGCAGGGGTGTAAAGAAAACACTTAG 364
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Genentech Inc. (US)
Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 473 from Patent W00140466. AX464340. AX464340.1 GI:21899186
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VERSION
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AUTHORS
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LOCUS
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SOURCE
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                                                                                                                                                              Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
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Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
                                                                                                                                                                                                                     Ashkenazi, A.J., Baker, K.P., Chan, B., Goddard, A., Godowski, P.J., Gurney, A.L., Hebert, C., Henzel, W., Kabakoff, R.C., Shelton, D.L., Tumas, D., Watanabe, C.K. and Wood, W.I.
Compositions and methods for the treatment of immune related diseases
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                                                                                                                                                                                                                                                                                                                             Patent: WO 0073452-A 82 07-DEC-2000;
                                                                                                                                                                                                                                                                                                                                                                                                     /organism="Homo sapiens"
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AX403489
            Sequence 82 from Patent WO0073452. AX055452
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Location/Qualifiers
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QY 181 GGCAAGGCATAGTTAAAGGACGGAATCTTGACTCAAGAGGGTTAATTCTTGGTGCTGAA 240 Db 272 GGCAAGGCATAGTTAAAGGACGGAATCTTGACTCAAGAGGGTTAATTCTTGGTGCTGAA 331 QY 241 GCCTGGGGCAGGGGTGTAAAGAAAACACTTAG 273 Db 332 GCCTGGGGCAGGGGTGTAAAGAAAACACTTAG 364 RESULT 5	AY358685 LOCUS AY358685 AY358685 DEFINITION Homo sapiens clone DNA60627 CLECSF12 (UNQ589) mRNA, complete cds. ACCESSION AY358685.1 GI:37182491 KEYWORDS FUL CDNA. SOURCE Homo sapiens (human) ORGANISM Homo sapiens Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo. REFERENCE AUTHORS Clark, H.F., Gurney, A.L., Abaya, E., Baker, K., Baldwin, D., Brush, J., Chen, J., Chow, B., Chui, C., Crowley, C., Currell, B., Deuel, B., Dowd, P., Eaton, D., Foster, J., Grimaldi, C., Gi, Q., Hass, P. E.,		AUTHORS Clark.H.F. AUTHORS Clark.H.F. JOURNAL Submitsed (01-AUG-2003) Department of Bioinformatics, Genentech, JOURNAL Submitted (01-AUG-2003) Department of Bioinformatics, JOURNAL SUBMITTED (11-AUG-2003) Department of Bioinformatics, JOURNAL SUB	CDS 1.054 7.10cus tag="UNQ589" 7.10cus tag="UNQ589" 7.10cus tag="UNQ589" 7.10cus tag="UNQ589" 7.10cus tag="UNG589" 7.10cus tag="UNG589" 7.10cus tag="UNG589" 7.10cus tag="UNG5812" 7.10cus tag="UNG581	Query Match 100.0%; Score 273; DB 9; Length 713; Best Local Similarity 100.0%; Pred. No. 2.6e-65; Indels 0; Gaps 0; Qy 1 ATGACATTTTTCTGTCACTATTATTATTGTTGGTATGTGAAGCTATTGGAGATCCAAT 60 Db 92 ATGACATTTTTCTGTCACTATTATTATTGTTGGTATGTGAAGCTATTTGGAGATCCAAT 151 Qy 61 TCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCAAGT 120 Db 152 TCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACACACATT Qy 121 CAACCCACACAATCATTAGAAGAAGAGTGTGACTCTTATCAAGAAATAAAGAAACACACTTTAGAAATAAAGAAACACACATTAGAAATAAAGAACACACATTAGAAAATAAAGAAAAAAAA

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1 ATGACAITITITCTGTCACIAITATIATIGTTGGTAIGTGAAGCTAITIGGAGAICCAAI
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       SEQUENCING READ COVERAGE:Sequencing is completed to a minimum standard of double strand coverage with a minimum of 2 clones and 2 reads with no ambiguities or 2 chemistries with a minimum of 2 clones and 3 reads with no ambiguities. If the sequence quality for a region does not meet this standard, it will be indicated in the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 note="Variable size of homopolymeric repeat in subclones"
function="unresolved homopolymeric run"
                                                                                                                                                                                                                                                                                                                                            CLONE LENGTH: This sequence does not necessarily represent the entire insert of this clone. Overlapping regions of clones are only sequenced and submitted once, so the sequence for the remainder of the insert may be found in the record for the adjacent clones. Overlapping clones are noted at the beginning and end of the Features listing.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Repeats are identified using RepeatMasker (A. Smit and P. Green, unpublished.) for Human and Mouse sequences.
Genes and Region of sequence similarity are identified by BLAST (Nuc. Acids Res. 25:3389-3402) similarity (expect < 1e-34) to the EST and cDNA sequences. Genes demonstrate at least two exons flanked by consensus splice sites that maintained sequence continuity across the splice junctions. Sequences that are not identical matches are annotated as similar.
                                                                                                                                                                                                   Submitted (29-MXY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA On May 29, 2002 this sequence version replaced gi:21206012. INFORMATION: http://www.hgsc.bcm.tmc.edu/ or email
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                STSs are identified using ePCR (Genome Res. 7:541-550) searches of a local database that includes entries from dbSTS, GDB, and local mapping efforts.
                                                                          Submitted (26-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, IX 77030, USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 QUALITY OF INDIVIDUAL BASES: This sequence meets stringent quality standards - estimated error rate less than 1 per 10,000 bases. Seports of lowest quality individual bases and measures of base quality and second of the metrics can be found quality are listed below. Description of the metrics can be found
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  http://gc.bcm.tmc.edu:8088/quality.info/genbank.annotation.html
        TX 77030, USA
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'rpt_family="(CA)n"

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complement(4558. 4629)
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/organism="Homo sapiens"
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/db_xref="taxon:9606"
/chromosome="12"
/clone="RP11-133114"
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/rpt_family="MLT1D"
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            Baylor Plaza, Houston,
                                                                                                                                                    (bases 1 to 165414)
                                (bases 1 to 165414)
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Direct Submission
                                             Worley, K.C.
Direct Submission
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AUTHORS
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NOTE: Estimated insert size may differ from sequence length (see http://www.hgsc.bcm.tmc.edu/docs/Genbank draft_data.html).
NOTE: This is a 'working draft' sequence. It currently consists of 16 contigs. The true order of the pieces is not known and their order in this sequence record is arbitrary. Gaps between the contigs are represented as runs of N, but the exact sizes of the gaps are unknown. This record will be updated with the finished sequence as soon as it is available and the accession number will
                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequencing vector: M13; L08821
Chemistry: Dye-primer Bodipy: 77% of reads
Chemistry: Dye-terminator Big Dye: 23% of reads
Assembly program: Phrap; version 0.990329
Consensus quality: 71594 bases at least Q40
Consensus quality: 110702 bases at least Q20
Estimated insert size: 229041; sum-of-contigs estimation
Batimated insert size: 31000; agarose-fp estimation
Quality coverage: 0.5x in Q20 bases; sqarose-fp estimation
Quality coverage: 0.9x in Q20 bases; sum-of-contigs estimation
                                                                                                       Submitted (05-FEB-1999) Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA On Sep 14, 2000 this sequence version replaced gi:10086352.
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111185: gap of unknown length
134782: contig of 23597 bp in length
134882: gap of unknown length
163654: contig of 28772 bp in length
163554: gap of unknown length
185050: contig of 21296 bp in length
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unknown length
of 8257 bp in length
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of 9652 bp in length
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                                                                                                                                                                                                                                                                                                                                                      Center project name: J-33
Center clone name: RP11-13C13, RP11-656E20
Sequencing vector: Plaemid; M77789
Sequencing vector: M13; L08821
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Contact: hgsc-help@bcm.tmc.edu
------- Project Information
                                                                                                                                                                                            ------ Genome Center
Center: Baylor College of Medicine
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238417: contig of 1171
238517: gap of unknown.
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                            2 (bases 1 to 240864)
Worley, K.C.
Direct Submission
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1 (Joses I LO 240864)

2 (Joses I LO 240864)

Alsbrooks, S.L., Amaratunge, H.C., Are, J.R., Banks, T., Barbaria, J., Benton, J., Brieva, M., Bloanker, E., Bonnin, D., Bouck, J., Benton, J., Brieva, M., Brown, E., Bryant, N.P., Bulay, C., Burch, P., Brieva, M., Brown, E., Bryant, N.P., Bulay, C., Burch, P., Burch, P., Burket, C., Burrell, K.L., Byrd, N.C., Carron, T.F., Carter, M., Cavazos, S.R., Chacko, J., Chen, C., Chen, R., Chen, Z., Chowdhry, I., Christopoulos, C., Cleveland, C.D., Cox, C., Coyle, M.D., Dathorne, S.R., David, R., Davila, M., Davis, C., Blaj, C., Brantz, P., Gabisi, A., Garler, P., Frantz, P., Gabisi, A., Garler, P., Frantz, P., Gabisi, A., Garler, P., Banks, C., Banks, C., Brants, C., Brants, C., Brants, C., Brants, C., Banks, C., Harris, K., Hart, M., Havlak, P., Hane, S., Hamilton, X., Harris, C., Harris, K., Hart, M., Havlak, P., Hane, J., Jackson, L., Jackson, L., Jackson, D., Tang, C., Liu, J., Li, Z., Lickarge, O., Lieu, C., Liu, J., Liu, W., Li, Z., Lickarge, O., Lieu, C., Liu, J., Liu, W., Mapua, P., Martina, E., Massey, R., Martinaz, E., Massey, E., Moser, M., Naul, D., Newtson, J., Newtson, J., Newtson, M., Morris, S., Moser, M., Roll, D., Newtson, E., Nowtson, M., Naul, M., Okwuonu, G., Oragunye, N., Oviedo, R., Pare, S., Parer, J., Palerisa, A., Tanerisa, A., Tanerisa, A., Tanerisa, A., Tanerisa, A., Tanerisa, K., Tanerisa, M., Phomas, R., Insan, Y., Vara, V., Villa, O., Villa, D., Villa, M., Charler, J., Lang, C., Taylor, C., Tayl
117208 CAACCCACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACA 117149
                                                                                                                                                                                                                                                                                                                                                         117148 GGCAAGGGCATAGTTAAAGGACGGAATCTTGACTCAAGAGGGTTAATTCTTGGTGCTGAA 117089
                                                                                                                                      117268 TCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCACAGT 117209
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Homo sapiens clone RP11-13C13, RP11-656E20, *** SEQUENCING IN AC006510 AC006510 AC006510, AC006510, AC006510, AC006514
                                                                                                                                                                                            121 CAACCCACAAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACA 180
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Thomas, S., Usmani, K., Vasquez, L., Vera, V., Villalon, D., Vinson, R., Wall, R., Wang, S., Ward-Moore, S., Warren, R., Washington, C., Watlington, S., Williams, G., Williamson, A., Wleczyk, R., Wooden, S., Worley, K., Wu, C., Wu, Y., Wu, Y.F., Zhou, J., Zorrilla, S., Nelson, D.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                    TCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCACAGT
                                                                                                                                                                                                                                                                                                      181 GGCAAGGGCATAGTTAAAGGACGGAATCTTGACTCAAGAGGGTTAATTCTTGGTGCTGAA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 117088 GCCTGGGCCAGGGCTGTAAAGAAAACACTTAG 117056
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Direct Submission
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AC006510/c
LOCUS
DEFINITION
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SOURCE ORGANISM

REFERENCE AUTHORS

TITLE

ACCESSION

VERSION KEYWORDS

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q ò qq à g ò q à g

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625 bp mRNA linear PRI 19-NOV-2001
Homo sapiens beta-glucan receptor isoform C (BGR) mRNA, complete
AP400597
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LAYLIGICITVILVIAVLYGTMALINRENSGSNTLENGYFLSRNKENHSQPTQSSLED
SVTPTKAVKTTGVLESPCPPRWIINERSCYLFSMSLNSWDGSKRQCWQLGSNLLKIDS
SNELISDQNHSYPKRPISKLCMDSRVSHL"
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                                                                                                                                                                                             GAAATAAAGAGAACCACAGTCAACCCACAATCATCTTTAGAAGACAGTGTGACTCCTA 160
                                                                                                                                                                                                                       310 GAAATAAAGAGAACCACAGTCAACCCACAAATCATCTTTAGAAGACAGTGTGACTCCTA 369
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /note="C-type lectin BGRC; similar to Mus musculus dectin
1; alternatively spliced"
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Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
1 (bases 1 to 625)
                                                                                                                                          41 AAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               43 GCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGA
                                                                             Gaps
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Willment, A.A., Gordon, S. and Brown, G.D.
Willment, A.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JUL-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
Location/Qualifiers
1. 625
/organism="Memo sapiens"
//mol_type="mRNA"
//db_xref="taxon:9606"
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Characterization of the human beta -glucan receptor and its
alternatively spliced isoforms
J. Biol. Chem. 276 (47), 43818-43823 (2001)
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                                      Length 881;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 tissue_type="peripheral blood leukocyte"
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100.0%; Pred. No. 7.5e-29;
ive 0; Mismatches 0;
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100.0%; Pred. No. 2.7e-28;
iive 0; Mismatches 0;
                                                                                                                                                                                                                                                                          CCAAAGCTGTCAAAACCACAGG 182
                                                                                                                                                                                                                                                                                                              370 ccaaagcrercaaaaccacagg 391
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            /chromosome="12"
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Best Local Similarity
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hes 142;
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AUTHORS
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Homo sapiens beta-glucan receptor isoform H (EGR) mRNA, complete
Cds, alternatively spliced.
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1 (bases 1 to 981)
Willment, J.A., Gordon, S. and Brown, G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
J. Biol. Chem. 276 (47), 43818-43823 (2001)
                                                                                                                                                                                                   211659 ATGACATTTTTCTGTCACTATTATTATTGTTGGTATGTGAAGCTATTTGGAGATCCAAT
                                                                                                                                                                                                                                                                                      211599 TCAGGAAGCAACACTTGGAGAATGGCTACTTTCTTATCAAGAAATAAAGAGAACCACAG
                                                                                                                                                                            1 ATGACATTTTTTCTGTCACTATTATTATTGTTGGTATGTGAAGCTATTTGGAGATCCAAT
                                                                                                                                                                                                                                                       TCAGGAAGCAACACATTGGAGAATGGCTACTTTC-TATCAAGAATAAAGAGAACCACAG
                                                                                                                                                                                                                                                                                                                                 T-CAACCCACAATCATCTTTAGAAGACAGTGTGACTCCTACC-AAAGCTGTCAAAACC
                                                                                                                                                                                                                                                                                                                                                                                                            A-CAGGCAAGGGCATAGTTAA-AGGACGGAATCTTGACTC-AAGAGGGTTAATTCTTGGT
                                                                                                                                      Gaps
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3RE, UK
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'codon_start=1
                                                                                             Length 240864;
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Willment, J.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JUL-2001) Sir William Dunn School
Oxford University, South Parks Road, Oxford OX1
Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       235 GCTGAAGCCTGGGGCAGGGTGTAAAGAAAAACACTTAG 273
                                                                                               DB 2;
                                                                                                                                    3;
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/clone="RP11-13C13, RP11-656E20"
                                                                                           Score 202.2; DB Pred. No. 1e-45; 0; Mismatches
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/chromosome="12"
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                                                                                               74.1%;
96.8%;
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                                                                                                               Best Local Similarity 96.8
Matches 270; Conservative
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AF400602
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LOCUS DEFINITION

RESULT 10 E21012

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KEYWORDS

SOURCE ORGANISM

AUTHORS TITLE

JOURNAL

COMMENT

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/translation="MEYHPDLENLDEDGYTQLHFDSQSNTRIAVVSEKGSCAASPPWR
LIAVILGILCLVILVIAVAGYLGTMAYWRSNSGSNTLENGYFLSRNKENHSQPTQSSLED
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SNBLGFIVKQVSQCPDNSFWIGLSRPQTEVWMIMEDGSTFSSNLFQIRTTATQENPSP
NCVWIHVSVIYDQLCSVPSYSICEKKPSM"
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1; alternatively spliced"
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 744)
                                                                                                                                                  Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Pathology,
                                                                                                                                                                                                      (bases 1 to 744)
Willment, J.A., Gordon, S. and Brown, G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
J. Biol. Chem. 276 (47), 43818-43823 (2001)
                                                                                                                                                                                                                                                                                                                                                                                                                                                             2 (bases 1 to 744)
Willment,J.A., Gordon,S. and Brown,G.D.
Direct Submissor
Submitted (18-JUD-2001) Sir William Dunn School of Patho
Oxford University, South Parks Road, Oxford OX1 3RE, UK
Location/Qualifiers
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Homo sapiens lectin-like receptor 1 (DECTIN1) mRNA,
AY026769
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protein id="AAL11711.1"
db_xref="GI:15986700"
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/organism="Homo sapiens"
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/db_xref="taxon:9606"
/chromosome="12"
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'gene="BGR"
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RS Shuji, H., Shoji, F., Kazunori, N. and Yasushi, A.

Novel membrane protein and DNA thereof

AL Patent: JP 1999001497-A 2 06-JAN-1999;

TAKEDA CHEM IND LTD
OS Unidentified

PN JP 1999001497-A/2

PD 06-JAN-1999

PF 13-JUN-1999 JP 1997156376

PR SHUJI HINUMA, SHOJI FUXUZUMI, KAZUNORI NISHI, YASUSHI ARAXA PC
COTKIA/47, A6IK38/00, A6IK38/00, A6IK38/00, A6IK39/395,

PC A6IK39/395,

PC A6IK48/00, COTKI6/28, CIZNI/21, CIZNI5/09, CIZPZ1/02.CIZO1/68. PC
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Homo sapiens beta-glucan receptor isoform A (BGR) mRNA, complete
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202 GCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGA
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/mol_type="genomic DNA"
/db xref="taxon:32644"
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Chem. 276 (47), 43818-43823 (2001)
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/db_xref="taxon:9606"
/chromosome="12"
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Homo sapiens (human)
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Hermanz-Falcon, P., Arce, I., Roda-Navarro, P. and Fernandez-Ruiz, E. Cloning of human DECTIN-1, a novel C-type lectin-like receptor gene expressed on dendritic cells [Immunogenetics 53 (4), 288-295 (2001)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   /traislation="MEYHPDIBNLDEDGYTQLHFDSQSNTRIAVVSEKGSCAASPPWR
LIAVILGILCLVILVIAVAVLGTWAJWRSNGSSNTLENGYFLSRNKENHSQPTQSSLED
SVPTKAVKTTGVLSSPCPPWJIYEKSCYLFSMSLNSWDGSKRQCWQLGSNLLKIDS
SNELGFIVKQVSSQPDNSFWTGLSRPQTEVPWLWEDGSTFSSNLFQIRTTATQENPSP
NCVWIHVSVIYDQLCSVPSYSICEKKFSM"
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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                                                                                                                                                           Submitted (01-FEB-2001) Biologia Molecular, Hospital de la Princesa, Diego de Leon 62, Madrid 28006, Spain 3 (bases 1 to 744)

Herranz-Falcon, P., Arce, I. and Fernandez-Ruiz, E. Direct Submission
Submitted (05-0CT-2001) Biologia Molecular, Hospital de la Princesa, Diego de Leon 62, Madrid 28006, Spain Sequence update by submitter on oct 5, 2001 this sequence version replaced gi:14278818.

Location/Qualifiers
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Herranz-Falcon, P., Arce, I. and Fernandez-Ruiz, E.
Direct Submission
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Isolated mammalian membrane protein gene and reagent relating
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Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
1 (bases 1 to 1018)
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JP 2002506645-A/3
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16-MAR-1999 JP 2000536856
17-MAR-1998 US 09/040111
LIOWEL CHALUS, AHN QUAN, ELIZABETH ESTHER MARY BATES, DANIEL M
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Isolated mammalian membrane protein gene and reagent relating Patent: JP 2002506645-A 3 05-MAR-2002;
SCHERING CORP
2 (bases 1 to 787)
Willment, J.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JUL-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
Location/Qualifiers
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cive 0; Mismatches 0; Indels
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Lipp,H.J., Mechtcheriakova,D., Schmid,J. and Sobanov,Y.
Likt variant, dino gene, lectin-like receptor gene, and proteins
encoded thereby
Patent: WO 02077216-A 7 03-OCT-2002;
Novartis AG (CH); Novartis Erfindungen Verwaltungsgesellschaft
C12N15/09, A61X31/70, A61X38/00, A61X39/395, A61X39/395, A61D35/00, A61D37/02, A61D43/00, C07X14/705, C07X16/28. C12N1/19 C12N1/19 C12N1/19
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25 37.6 13.8 595 9 26 37.4 13.7 519 7 27 37 13.6 150 7 29 37 13.6 866 8 30 37 13.6 866 8 31 37 13.6 868 8 31 37 13.6 868 8 31 37 13.6 808 8 33 37 13.6 808 8	35 37 13 6 912 8 8 8 36 8 13 5 7 13 6 912 8 8 8 36 6 13 4 5 74 8 8 6 8 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 13 6 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	36 13.2 717 5 3.6 13.2 717 5 3.6 13.2 73.4 5 5 3.6 13.2 73.4 5 5 3.6 13.2 73.4 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	
GenCore version 5.1.6 Copyright (c) 1993 - 2004 Compugen Ltd. OM nucleic - nucleic search, using sw model Run on: December 1, 2004, 23:06:11; Search time 1366.11 Seconds (without alignments) 7282.038 willion cell undates/sec	aaagaaaaacacttag 273	Database: EST:* 1: gb_est1:* 2: gb_est2:* 3: gb_tc:* 4: gb_est3:* 5: gb_est4:* 6: gb_est5:* 7: gb_est5:* 7: gb_est5:* 8: gb_gss1:* 9: gb_gss1:* 9: gb_gss1:* 9: gb_gss2:* Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution. SUMMARIES	No. Score Match Length DB ID Description

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VERSION KEYWORDS SOURCE ACCESSION

RESULT 2 CB958894

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Eukaryotza, Metazoa, Chordata, Craniata, Vertebrata; Euteleostomi;
Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.

1 (bases 1 to 659)
Dias Neto, E., Garcia Correa, R., Verjovski-Almeida, S., Briones, M.R., Nagai, M.A., da Silva, W. Jr., Zago, M.A., Bordin, S., Costa, F.F., Brunstein, G.H., Carvalho, A.F., Matsukuma, A., Baia, G.S., Simpson, D.H., Brunstein, A., deoliveira, P.S., Bucher, P., Jongeneel, C.V., O'Harre, M.J., Soares, F., Brentani, R.R., Reis, L.F., de Souza, S.J. and
                                                                                                                                               BI018962 659 bp mRNA linear EST 14-JUN-2001 IL3-MT0267-050101-437-A02 MT0267 Homo sapiens CDNA, mRNA sequence.
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Fax: +55-11-2707001
Email: asimpson@ludwig.org.br
This sequence was derived from the FAPESP/LICR Human Cancer Genome Project. This entry can be seen in the following URL (http://www.ludwig.org.br/scripts/gethtml2.pl?tl=IL3&t2=IL3-MT0267-0550101-437-A02&t3=2001-01-05&t4=1)
Seq primer: puc 18 forward
High quality sequence stop: 639.
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Laboratory of Cancer Genetics
Ludwig Institute for Cancer Research
Rua Prof. Antonio Prudente 109, 4 andar, 01509-010, Sao Paulo-SP,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Shotgun sequencing of the human transcriptome with ORF expressed
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Proc. Natl. Acad. Sci. U.S.A. 97 (7), 3491-3496 (2000)
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Pred. No. 3.1e-29;
0; Mismatches 1; Indels 0;
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Sfil (ggccattatggcc); Site_2: Sfil (ggccgcctcggcc);
Library is oligo-dT primed and directionally cloned. cDNA
was prepared from a glandular pool of tissues from thyoid,
parathyroid, adreal, cortex and pineal gland. 5' and 3'
adaptors were used in cloning as follows: 5' adaptor
sequence: 5'-CACGCCGATTATGGCC-3' and 3' adaptor sequence:
5'-ATTCTAGAGGCCGAGGCGCGACATG-dT(310) Bn.3' (where B = A,
C, or G and N = A, C, G, or T). Average insert size 1.38
kb (range 0.60-3:5 kb): 15/15 colonies contained inserts
by PCR. This library was enriched for full-length clones
and was constructed by Clontech Laboratories (Palo Alto,
CA). Note: this is a NIH_MGC Library."
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Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 800)
WHH-MGC http://mgc.nci.nih.gov/.
National Institutes of Health, Mammalian Gene Collection (MGC)
Unpublished (1999)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CDNA Library Preparation: CLONTECH Laboratories, Inc. CDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL) DNA Sequencing by: Apencourt Bioscience Corporation Clone distribution: MGC clone distribution information can http://image.llnl.gov.e. column: 19 Plate: NDCM151 row: e column: 19 Plate: MCM151 row: e column: 19 High quality sequence stop: 488.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Contact: Robert Strausberg, Ph.D.
Email: cgapbs-r@mail.nih.gov
Tissue Procurement: Dr. Michael Brownstein and Dr. Miklos
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ;
                                                                                                                                                                                   AGENCOURT 13785021 NIH MGC 184 Homo sapiens cDNA clone IMAGE:30352338 5', mRNĀ sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 800;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 6;
240 AGCCTGGGCCAGGGGTGTAAAGAAAACACTTAG 273
                              271 AGCCTGGGGCAGGGTGTAAAGAAAACACTTAG 238
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1.6e-40;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ilarity 98.4%; Score 177.2; 98.4%; Pred. No. 1.6e Conservative 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       organism="Homo sapiens"
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1. .800
                                                                                                                                                                                                                                                                                              CB958894.1 GI:30215010
                                                                                                                                                                                                                                                                                                                                                      Homo sapiens (human)
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Matches 179;
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AV721179 GV SAPPENS CON 673 bp mRNA linear EST 16-OCT-2000 AV721179 HTB Homo sapiens CDNA clone HTBAKF11 5', mRNA sequence.
                                                                                                                                                                                                                                                                                                                      Homo sapiens

Bukaryota, Merazoa; Chordata; Craniata; Vertebrata; Buteleostomi;

Bukaryota; Dutheria; Primates; Catarrhini; Hominidae; Homo.

(bases 1 to 673)

(bu,Y., Peng,Y., Song,H., Huang,Q., Yang,Y., Gao,G., Xiao,H., Xu,X.,

(bu,Y., Peng,Y., Song,H., Huang,Q., Yang,Y., Gao,G., Xiao,H., Xu,X.,

Xu,S., Gu,W., Tu,Y., Jia,J., Gu,J., Gao,X., Cheng,Z., Xu,Z., Zeng,L.,

Xu,S., Gu,W., Tu,Y., Jia,J., Fu,G., Ren,S., Zhong,M., Lu,G., Hu,R.,

Chen,J., Chen,Z. and Han,Z.

Unpublished (2000)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              EST 25-MAR-2003
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   123 ACCCACACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACAGG 182
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1 (Dases 1 to 582)
Smith,T.P.L., Roberts,A.J., Echternkamp,S.E., Chitko-McKown,C.G.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          /clone_lib="HTB"
/note="Vector: pBluescript sk(-); Site_1: EcoRI; Site_2:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Contact: Zeguang Han
Contact: Zeguang Han
Chinese National Human Genome Center at Shanghai
351 Guo Shoujing Road, Zhangjiang Hi-Tech Park, Pudong, Shanghai
201203, P. R. China
Tel: 86-21-50801919(ex.45)
Fax: 86-21-50801922
Email: hanzg@chgc.sh.cn
This clone is available at CHGC in Shanghai.
Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      1 AGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAGAAATAAAGAGAACCACAGTCA
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Pred. No. 6.9e-24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              CB420818 582 bp mRNA linear 593806 MARC 6BOV Bos taurus cDNA 5', mRNA sequence.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             tissue type="Hypothalamus"
|dev stage="Adult"
|lab_host="SOLR"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         100.0%; Pred. NC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             /organism="Homo sapiens"
/mol_type="mRNA"
/db_xref="taxon:9606"
/clone="HTBAKF11"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CB420818
CB420818.1 GI:29187264
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243 CTGGGGCAGGG 253
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CB420818
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/note="Organ: Kidney/Brain, Vector: pBACe3.6; Site 1:
EcoR; Site 2: EcoR; Female C57BL/6J mouse kidney and/or
brain genomic DNA was isolated and partially digested
with a combination of EcoRI and EcoRI Methylase. Size
selected DNA was cloned into the pBACe3.6 vector at the
EcoRI sites. The ligation products were transformed into
DH10B electrocompetent cells (BRL Life Technologies). "
                           AZ121459 663 bp DNA linear GSS 12-MAY-2000 RPCI-23-30A15.TV RPCI-23-30A15.TV RPCI-23-30A15.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             190 GAAATCCAGAGGAGAAAAACAACTTCCTATCAAGAATAAAGAGAACCACA--AGCCCA 134
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             CACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACAGGCAAGG 187
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              GCATAGITAAAGGA-----CGGAATCITGACTCAAGAGGGTTAATTCTTGGTGCTGAAGC 242
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Email: szhaogtigr.org

Email: szhaogtigr.org

Clones are derived from the mouse BAC library RPCI-23. For BAC

library availability, please contact Pieter de Jong

(pieter@dejong.med.buffalo.edu). Clones may be purchased from

BACPAC Resources (http://bacpac.med.buffalo.edu/orderingframe.htm)

or from Resea ch Genetics (info@resgen.com). BAC end page:

http://www.tigr.org/tdb/bac_ends/mouse/bac_end_intro.html

Seq primer: T7
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                                                                                                                                                                                                                                                                                       Zhao, S., Nierman, W., Feldblyum, T., Malek, J., Shatsman, S., Akinret, B., Lewins, M., Mcgann, S., Tsegaye, G., Geer, K., Krol, M., de Jong, P. and Fraser, C.M.
                                                                                                                                                                                                         Mus musculus
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Butheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 663)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 67
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Pred. No. 4.6e-24;
0; Mismatches 59; Indels 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                        Contact: Shaying Zhao
Department of Eukaryotic Genomics
The Institute for Genomic Research
712 Medical Center Dr., Rockville, MD 20850, USA
71: 301 838 0200
Fax: 301 838 0208
                                                                                                                                                                                                                                                                                                                                                                    Mouse BAC End Sequences from Library RPCI-23 Unpublished (1999) Other_GSSs: RPCI-23-30A15.TJ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /organism="Mus musculus"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     /mol_type="genomic DNA"
/strain="C57BL/6J"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     /db_xref="taxon:10090"
/clone="RPCI-23-30A15"
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                                                                                                                                                                               Mus musculus (house mouse)
                                                                                   genomic survey sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        /sex="Female"
                                                                                                                          AZ121459.1 GI:7788387
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ilarity 73.3%;
Conservative (
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BY547544
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                                                                      USDA, ARS, US Meat Animal Research Center
PO Box 166, Clay Center, NE 6933-0166, USA
Tel: 402 762 4366
Fax: 402 762 4360
Email: smith@email.marc.usda.gov
Single pass sequencing. Bases called with phred v0.020425.c and
trimmed with the aid of the trim_alt option. Vector identified with
Plate: FQY8014 row: B column: 7
Seq primer: GTAARAGACTCACTATAGGG.
  A second set of bovine ESTs from pooled-tissue normalized libraries Unpublished (2003)
Contact: Smith TPL
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       BI107684 855 bp mRNA linear EST 26-JUN-2001
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     43 GCTATITGGAGATCCAATICAGGAAGCAACACATIGGAGAATGGCTACTITCTATCAAGA 102
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            244 GGTATTTGGAGATCCAGTTCAGGGAACAACCTGTTGAAGAGTGACAGCTTTCCATCAAGA 303
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      103 AATAAAGAAACCACAGTCAACCCACACAATCATCTTTAGAAGACAGTGTGACTTCCTACC 162
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
1 (bases 1 to 855)
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/clone_lib="MARC 6BOV"
/note="Vector: pcDNa3.1; Site_1: EcoRI; Site_2: NotI;
Libzary made with RNA pooled from multiple tissues
including liver, lung, hypothalamus, pituitary, and
placenta/endometrium."
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Contact: Robert Strausberg, Ph.D.
Email: cgapbs-r@mail.nih.gov
Tissue Procurement: Gilbert Smith, Ph.D.
cDNA Library Preparation: Life Technologies, Inc.
cDNA Library Arrayed by: The I.M.A.G.B. Consortium (LLNL)
DNA Sequencing by: Incyte Genomics, Inc.
Clone distribution: MGC clone distribution information can be
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  NIH-MGC http://mgc.nci.nih.gov/.
National Institutes of Health, Mammalian Gene Collection (MGC)
Unpublished (1999)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  .;
0
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              found through the I.M.A.G.E. Consortium/LLNL at: http://image.llnl.gov
Plate: LLAM1101 row: j column: 08
High quality sequence start: 27
High quality sequence stop: 825.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      38.4%; Score 104.8; DB 6 ilarity 84.3%; Pred. No. 1.8e-19; Conservative 0; Mismatches 22
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                                                                                                                                                                                                                                                                                          Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         163 AAAGCTGTCAAAACCACAGG 182
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Wray, J.E. and Keele, J.W.
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Matches 118; Conserv
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mas musalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Musses I (bases I to 363)

RS Okazaki, Y., Furuno, M., Kasukawa, T., Adachi, J., Bono, H., Kondo, S., Nikaido, I., Osato, N., Saito, R., Suzuki, H., Yamanaka, I., Sthosawa, H., Yaqi, K., Tomaru, Y., Hasegawa, Y., Nogami, A., Schonbach, C., Gojobori, T., Baldarelli, R., Hill, D.P., Bult, C., Grobani, I.E., Cousins, S., Dalla, E., Dragani, T., Hume, D.A., Quackenbush, J., Schrimi, L.M., Kanapin, A., Matsuda, H., Chothia, C., Corbani, L.E., Cousins, S., Dalla, E., Dragani, T.A., Fletcher, C.F., Forrest, A., Frazer, K.S., Gaasterland, T., Gariboldi, M., Gissi, C., Godzik, A., Gough, J., Grimmond, S., Gavich, M., Hirokawa, N., Jackson, I.J., Jarvis, E.D., Kanai, A., Kurochkin, I.V., Lee, Y., Lenhard, B., Lyons, P.A., Maglott, D.R., Wurochkin, I.V., Lee, Y., Lenhard, B., Lyons, P.A., Maglott, D.R., Numata, K., Okido, T., McKenzie, L., Miki, H., Nagashima, T., Numata, K., Okido, T., Pertea, G., Pesole, G., Reed, D.G., Read, G., Pesole, G., Semple, C.A., Satole, R., Vananabe, Y., Saltenaka, Y., Takenaka, Y., Taylor, M.S., Teasdale, R.D., Tomita, M., Verardo, R., Wanner, A., Cavolan, M., Zhu, Y., Zhu, Yang, Y., Yasunish, Y., Sasaki, M., Jandan, K., Itoh, M., Kagawa, T., Konno, H., Nakawa, T., Shibata, K., Shibata, K., Sasaki, K., Sasaki, K., Sasaki, K., Sasaki, K., Shibata, K., Shibata, K., Shibata, K., Shibagawa, R., Shibada, Y., Shibata, K., Shibata,
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               42 AGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCTACTTTCTATCAAG 101
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  310 AGCATTATGGAGACACAATTCAGGGAGAATTCCAGAAGAGAAAGACAACTTCCCATCAAG 369
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           102 AAATAAAGAGAACCACAGTCAACCCACAATCATCTTTAGAAGACAGTGTGACTTCTTAC 161
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Site_2: Not1; Cloned unidirectionally. Primer: Oligo dT.
Library constructed by Life Technologies. Investigator
providing samples: Gilbert Smith, NIH"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 370 AAATAAAGAGAAACCACA---AGCCCACAGAATCATTTAGATGAGAAGGTGGCTCCCTC 426
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                3;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        27.0%; Score 73.8; DB 4; 75.2%; Pred. No. 1.9e-10; iive 0; Mismatches 32;
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Location/Qualifiers
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       162 CAAAGCTGTCAAAACCACAGG 182
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                                                                                                                                                    type="mRNA"
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Best Local Similarity
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JOURNAL MEDLINE PUBMED

COMMENT

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Nikaido, I., Osato, N., Saito, R., Adachi, J., Bono, H., Kondo, S., Nikaido, I., Osato, N., Saito, R., Siandarelli, R., Hill, D. P., Bult, C., Golobori, T., Daldarelli, R., Hill, D. P., Bult, C., Hayosawa, M., Osato, N., Basegawa, Y., Nogami, A., Riyosawa, H., Wagi, K., Tomaru, Y., Hasegawa, Y., Nogami, A., Batalov, S., Beisel, K., Baldarelli, R., Hill, D. P., Bult, C., Gondoni, J., B., Cousins, S., Dalla, E., Dragani, T.A., Fletcher, C. Corbani, L.E., Cousins, S., Dalla, E., Dragani, T.A., Fletcher, C. Corbani, L.E., Cousins, S., Gasterland, T. Kanai, H., Kawai, K., Grimmond, S., Gustinocich, S., Hirokawa, N., Jackson, I.J., Jarvis, E.D., Kanai, A., Kawaii, H., Kawasawa, Y., Kedzierski, R.M., King, B.L., Konagaya, A., Karochini, I.V. Lee, Y., Lehard, B., Lyons, P.A., Maglott, D.R., Kurochini, I., Marchionni, L., McKenzie, L., Miki, H., Nagashima, T., Nimata, K., Okido, T., Pertea, G., Pectrovsky, N., Pillai, R., Ponttius, J.U., Qi, D., Ramachandran, S., Petrovsky, N., Pillai, R., Ponttius, J.U., Qi, D., Ramachandran, S., Ravasi, T., Wagner, L., Wanlestedt, C., Wang, Y., Watanabe, Y., Wagner, L., Wanlestedt, C., Wang, Y., Watanabe, Y., Wang, L., Wanlestedt, C., Wang, Y., Watanabe, Y., Wang, L., Yan, Z., Zavolan, M., Zhu, Y., Zimmer, A., Carninci, P., Hayatsu, M., Hirozane-Kishikawa, T., Waki, K., Kawai, J., Aizawa, K., Sakazume, N., Sato, K., Shiraki, T., Waki, K., Kawai, J., Aizawa, K., Itoh, M., Kagawa, I., Miyazaki, A., Sasaki, D., Sasaki, D., Shibata, R., Rogers, J., Birray, B., Hara, A., Yashinagawa, A., Yasunishi, A., Yoshino, M., Waterston, R., Lander, E.S., Analysis of the mouse transcriptome based on functional annotation of 60,770 full-length cDNas
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Laboratory for Genome Exploration Research Group, RIKEN Genomic Sciences Center(GSC), Yokohama Institute
Sciences Center(GSC), Yokohama Institute
The Institute of Physical and Chemical Research (RIKEN)
1-7-22 Subhiro-cho, Tsurumi-ku, Yokohama, Kanagawa 230-0045, Japan Tel: 81-45-503-9216
Email: genome-reseascs.riken.jp, URL:http://genome.gsc.riken.jp/
Aizawa,K., Akimura,T., Arakawa,T., Carninci,P., Fukuda,S.,
Hirozane,T., Imotani,K., Ishii,Y., Itoh,M., Kawai,J., Konno,H.,
Miyazaki,A., Murata,M., Nakamura,M., Nomura,K., Numazaki,R.,
Ohno,M., Sakai,K., Sakai,K., Sasaki,D., Sato,K., Shibata,K.,
Shiraki,Y., Tagami,M., Waki,K., Watahiki,A., Muramatsu,M. and
Hayashizaki,Y. Direct Submission
Computentional Analysis of Pull-Length Mouse cDNAs Compared with
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human Genome Sequences Mamm. Genome. 12, 673-677 (2001)
Normalization and subtraction of cap-trapper-selected cDNAs to prepare full-length cDNA libraries for rapid discovery of new genes. Genome Res. 10 (10), 1617-1630 (2000)
RIKEN integrated sequence analysis (RISA) system--384-format sequencing pipeline with 384 multicapillary sequencer. Genome Res. 10 (11), 155-1771 (2000)
Computer-based methods for the mouse full-length cDNA encyclopedia: real-time sequence clustering for construction of a nonredundant cDNA library. Genome Res. 11 (2), 281-289 (2001) cDNA library was prepared and sequenced in Mouse Genome Encyclopedia Project of Genome Exploration Research Group in Riken Genomic Sciences Center and Genome Science Laboratory in RIKEN.
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Tissues were provided by Dr. John Todd (Dept. of Medical Genetics
Wellcome Trust Centre for Molecular Mechanisms in Disease Wellcome
Trust/MRC building Addenbrookes Hospital Cambridge) whose
                                                                                                                                                                                                                             Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
                                                                                                                                                                                       Chordata; Craniata; Vertebrata; Buteleostomi;
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Okazaki,Y., Furuno,M., Kasukawa,T., Adachi,J., Bono,H., Kondo,S.,
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                                                                                                            Mus musculus (house mouse)
Mus musculus
Eukaryota; Metazoa; Chorda
                                 GI:26871045
BY536666
BY536666.1
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Normalization and subtraction of cap-trapper-selected cDNAs to prepare full-length cDNA libraries for rapid discovery of new genes. Genome Res. 10 (10), 1677-1630 (2000)
RIKEN integrated sequence analysis (RISA) system-184-format sequencing pipeline with At multicapillary sequencer. Genome Res. 10 (11), 1757-1771 (2000)
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                                                                                                                                                                                                                                                                       Laboratory for Genome Exploration Research Group, RIKEN Genomic Sciences Center(GSC), Yokohama Institute
The Institute of Physical and Chemical Research (RIKEN)
1-7-22 Suehiro-cho, Tsurumi-ku, Yokohama, Kanagawa 230-0045, Japan Tel: 81-45-503-9226
Fax: 81-45-503-9216
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Computer-based methods for the mouse full-length cDNA encyclopedia: real-time sequence clustering for construction of a nonredundant cDNA library. Genome 8.11 (2), 281-289 (2001) cDNA library was prepared and sequenced in Mouse Genome Encyclopedia Project of Genome Exploration Research Group in Riken Genomic Sciences Center and Genome Science Laboratory in RIKEN. Division of Experimental Animal Research in Riken contributed to
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      prepare mouse Lissues.
Tissues were provided by Dr. John Todd (Dept. of Medical Genetics Wellcome Trust Centre for Molecular Mechanisms in Disease Wellcome Trust (MRC building Addenbrookes Hospital Cambridge) whose assistance we gratefully acknowledge.
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                                     Analysis of the mouse transcriptome based on functional annotation of 60,770 full-length cDNAs Nature 420, 563-573 (2002)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        67
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/clone_lib="RIKEN full-length enriched, B6-derived CD11
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Email: genome-res@gsc.riken.jp, URL:http://genome.gsc.riken.jp/
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Pred. No. 0.0018;
        Birney, E. and Hayashizaki, Y.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         0; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         /organism="Mus musculus"
/mol type="mRNA"
/strain="C57BL/6J"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              /db_xref="taxon:10090"
/clone="F730119008"
                                                                                                                                                                                                                                              Contact: Yoshihide Hayashizaki
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    .ve dendritic cells"
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72.7%;
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Query Match

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FEATURES

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Genome Research Group

National Institute of Agrobiological Sciences
Owashi 1-2, Tsukuba, Ibaraki 305-8634, Japan
Email: kmita@nias.affrc.go.jp
method:uni-directional, sequence direction:sequenced from T3 primer
                                                                                                                                                                                                                                       BP117361 e-- Bombyx mori cDNA clone ce--0217, mRNA sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AV882594 Satoh unpublished cDNA library, larva Ciona intestinalis cDNA clona cilv29904 3', mRNA sequence.
        124 CCCACACAATCATCTTTAGAAGACAGTGTGACTCCTACCAAAGCTGTCAAAACCACAGGC 183
                                             305 rėcaakirirerarearaarerearareakiesis 146
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  157 CCTACCAAAGCTGTCAAAACCACAGGGCAAAGGGCATAGTTAAAGGACGGAATCTTGACTCA 216
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Phlebobranchia; Cionidae; Ciona.
1 (bases 1 to 516)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     236 GTGAAGAAATCTTTAAAATTCACAGAAACTACGACAAATAATCCACTGAAACTAGAGATA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   97 TCAAGAAATAAAGAGAACCACAGTCAACCCACACAATCATCTTTAGAAGACAGTGTGACT
                                                                                                                                                                                                                                                                                                                                                                                   Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota; Neoptera; Endopterygota; Lepidoptera; Glossata; Ditrysia; 1 (bases 1 to 450) Mita, K., Morimyo, M., Shimada, T., Okano, K. and Maeda, S. Bombyx mori cDNA (Mita, K. 2003) Unpublished (2003)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          296 AGCGATICTAGIGITAGIACIAACGGCICTCCCACTAAACIAAGGCTATACACAT 350
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       /tissue type="compound eye"
/clone_Tib="ce--"
/note="mixed stages from 5th instar larva to pupa"
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/mol_type="mRNA"
/db_xref="taxon:7091"
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Bombyx mori
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                                                                                         184 AAGGCATAGTTAAAG 199
                                                                                                                                245 AAGTGCAGATACAAAG 230
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/clone_lib="RIKEN full-length enriched, NOD-derived CD11c
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1 (100 822)
Satou,Y., Shin-i,T., Kohara,Y. and Satoh,N.
Expressed genes in Ciona intestinalis (2002c)
Unpublished (2002)
Contact: Nori Satoh
Department of Zoology
                                                                                                                                                                                                                                                                                                                                                                                                                                        332 TTTTTCTGTCACTATTGCTGTTCGTGACGTATGAAGCATTTTGGCGACACAATTCAGGGA 391
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               /organism="ciona intestinalis"
/mol type="mRNA"
/db_xef="teaxon:719"
/clone="roid037h12"
/tissue_type="gonad"
/clone_lib="Nori Satoh unpublished cDNA library, gonad"
assistance we gratefully acknowledge.
Please visit our web site (http://genome.gsc.riken.go.jp) for
further details.
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Pred. No. 0.0019;
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Sakyo-ku, Kyoto, Kyoto 606-8502, Japan
Tel: 81-75-753-4081
Fax: 81-75-705-1113
Email: satch@ascidian.zool.kyoto-u.ac.jp.
Location/Qualifiers
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                                                                                                    /organism="Mus musculus"
/mol_type="mRNA"
/strain="NOD"
                                                                                                                                                                 /db_xref="taxon:10090"
/clone="F630311P20"
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Ciona intestinalis
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BW404792 Yutaka Satou unpublished cDNA library, embryo whole animal Ciona intestinalis cDNA clone ciem846h18 3', mRNA sequence.
BW404792 GI:47820620
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/tissue_type="whole animal"
/dev stage="embryo"
/clone_lib="Yutaka Satou unpublished cDNA library, embryo
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Bukaryota; Metazoa; Chordata; Urochordata; Ascidiacea; Enterogona; Phlebobranchia; Cionidae; Ciona.

1 (bases 1 to 703)
Satou, Y., Shin-i, T., Kohara, Y. and Satoh, N.
Expressed genes in Ciona intestinalis (2004)
Unpublished (2004)
Contact: Yutaka Satou
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Tel: 81-75-75-4095
Fax: 81-75-705-1113
Email: yutakaascidian.zool.kyoto-u.ac.jp.
Location/Qualifiers
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        Satoh, N., Satou, Y., Kohara, Y. and Shin-i, T. Expressed genes in Ciona intestinalis Unpublished (2000) Contact: Nori Satoh Department of Zoology Kyoto University Sakyo-ku, Kyoto, Kyoto 606-8502, Japan Tel: 81-75-753-4081
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Contact: Nori Satoh
Department of Zoology
Kyoto University
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5.1.6 Compugen Ltd.	; Search time 3773.98 Seconds (without alignments) 8934.210 Million cell updates/sec	tgttgccaaggaaaaaaa 713		gnes	ers: 9053458						4000	b predicted by chance to make a score of the result being printed, total score distribution.	Description	AX055452 Sequence AX055452 Sequence	AX464340 Sequence AX458688 Home sapi	ACC4224 NONE SAPI ACC06510 Home sapi AC138620 Mus muscu	AC112033 Rattus no AF400602 Homo sapi	AF400597 Homo sapi E21012 Novel membr NEVACES Homo sari	AYOOOO HOMO SAFI AYOOO HOMO SADI AF400601 Homo SADI	BD136999 Isolated AX664609 Sequence	AJ3123/3 Homo Sapı CQ779237 Sequence
GenCore version 5 Copyright (c) 1993 - 2004 CO	on: December 1, 2004, 23:10:21	Title: US-09-989-293A-376 Perfect score: 713 Sequence: 1 aatatatcatctatca	Scoring table: IDENTITY_NUC Gapop 10.0 , Gapext 1.0	Searched: 4526729 segs, 23644849745 residue	Total number of hits satisfying chosen parameters Minimum DB seq length: 0	sed rengin: ssing: Minimu Maximu Listin	Database : GenEmbl:* 1: gb ba:* 2: gb_htg:*	3: gb_in:* 4: gb_om:* 5: gb_ov:* 6: gb_pat:*	7: da da 7 8: da da 1:* 9: da da :*	10: gb_ro:* 11: gb_sts:* 12: gb_sy:* 13: gb_un:*	14: gb_vi:*	resurt 1 to th of the	\$ Result Query No. Score Match Length DB ID	713 100.0 713 6	4 713 100.0 5 713 100.0	705:6 99:0 165414 9 590.2 82.8 240864 2 186 26.1 221471 2	147.6 20.7 273867 2 10 142 19.9 881 9	1 140 19.6 625 9 2 140 19.6 741 6	140 19.6 744 9 140 19.6 744 9 140 19.6 787 9	6 140 19.6 1018 6 7 140 19.6 1153 6	8 140 19.6 1153 9 9 140 19.6 1606 6

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Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
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                 GTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACAACAATTGGAGAATGGCTA
                                                         CTTTCTATCAAGAAATAAAGAGAACCACAGTCAACCCACACAATCATCTTAGAAGACAG
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100.0%; Pred. No. 1.5e-135;
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Sequence 473 from Patent W00140466.
AX464340
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Bukaryota; Metazoa; Chordata; Catarrhini; Hominidae; Homo.

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Clark, H.F., Gurney, A.L., Abaya, E., Baker, K., Baldwin, D., Brush, J.,

Chen, J., Chow, B., Chhi, C., Crowley, C., Currell, B., Deuel, B.,

Dowd, P., Eaton, D., Foster, J., Grimaldi, C., Gu, Q., Hass, P.E.,

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Stinson, J., Vangler, A., Vandlen, R., Watanabe, C., Mieand, D., Woods, K.,

Xie, M.H., Yansura, D., Yi, S., Yu, G., Yuan, J., Zhang, M., Zhang, Z.,

Goddard, A., Wood, M.I. and Godowski, P.

The Secreted Protein Discovery Initiative (SPDI), a Large-Scale

Effort to Identify Novel Human Secreted and Transmembrane Proteins:

A Bioinformatics Assessment

M. Genome Res. 13 (10), 2265-2270 (2003)
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Homo sapiens clone DNA60627 CLECSF12 (UNQ589) mRNA, complete cds.
AY3S8685
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               GTATCCCTGTCATATATACAATAAGGTGAAATTATAAGTACCCTATGCAGTTGGGTGGAC
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                                                                                                                                                             121 GTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCTA
                                                                                                                                                                                                                                                                                                                                   TGACTCAAGAGGGTTAATTCTTGGTGCTGAAGCCTGGGGCAGGGGTGTAAAGAAAAACAC
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TGTGACTCCTACCAAAGCTGTCAAAACCACAGGCCAAGGGCATAGTTAAAGGACGGAATCT
                                                                                                                                                                                                                                                                                 241 TGTGACTCCTACCAAAGCTGTCAAAACCACAGGGCAAGGGCATAGTTAAAGGACGGAATCT
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                                                      TITIGGGATITITAATITICAAACACAGCAGAATGACATITITITICIGICACTATIATIT
                                                                                                                         GTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAAATGGCTA
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Mammallar, Eutherlay, Frinates; Catarrhin; Honinidae; Homo.

RS MurryD. M., Admas. C., Adio-Oducia B., All-osman, F. R., Allen, C.,

Alabrooks, S. L., Amartunge, H. C., Are, J. R., Ayele, M., Banks, T.,

Barbaria, J., Benton, J., Bimage K., Blankenburg, K., Bonnin, D.,

Bunky, C., Burch, P., Burket, C., Burrell, K. L., Byrd, N. C.,

Carron, T. R., Catfer, M., Cavacos, R., Chocko, J., Chavez, D.,

Chen, G., Chen, R., Chen, Z., Chiu, D., Dathorne, S. R., David, N. C.,

Clarch, T. C., Cox, C., Coyle, M. D., Dathorne, S. R., David, R.,

Delaney, K. R., Delgado, D., Denn, A. L., Ding, Y., Dinh, H. H.,

Delaney, K. R., Delgado, D., Denn, A. L., Ding, Y., Dinh, H. H.,

Douthwaite, K. J., Davis, C., Davyle, Carroll, L., Dederich, D. A.,

Barnhart, C., Eddar, M., Gao, J., Carroll, L., Dederich, D. A.,

Gill, R., Gorrell, J. H., Gao, J., Garcia, A., Garner, T., Garza, N.,

Frantz, P., Gabisi, A., Gao, J., Harris, K., Hart, M., Havlik, P.,

Hamilton, K., Han, J. Harris, C., Harris, K., Hart, M., Hale, S.,

Handloway, C., Han, J., Harris, C., Harris, K., Hart, M., Hale, S.,

Hum, J., Diliver, S., Joudah, S., Machen, B., Jia, Y.,

Joliver, M., Miller, M., Martindale, A., Marthez, B., Jia, Y.,

Leals, B., Lee, L., Lewis, L. C., Lewis, L. C., Lewis, L. L., Lothargo, N.,

Marniney, B., Morting, B., Marting, B., Jia, Y.,

Lieu, C., Liu, J., Liu, W., Loolesged, H., Lozado, R. J., Lu, X.,

Marniney, R., Molter, R., Luna, R., Ma, J., Maheshwari, M., Mapua, P.,

Marniney, R., Molter, R., Luna, R., Ma, J., Maheshwari, M., Mapua, P.,

Marniney, R., Molter, R., Luna, R., Ma, J., Maheshwari, M., Mayue, N.,

Mayben, N., Mickerson, J., Wertson, J., Wither, S., Saoer, J.,

Scherk, J., Willer, A., Jonespan, M., Maheshwari, M., Muyen, N.,

Nelyen, N., Mickerson, J., Werson, J., Wather, S., Saoer, J.,

Scherk, J., Wayene, R., Wayerson, J., Wather, S., Saoer, J.,

Scherk, J., Wayene, N., Willer, M., Wather, M., Maheshwari, M., Wayen, N.,

Scherk, J., Wayene, N., Wayerek, A., Tang, H.,

Sutek, A., Wayerek, A., Tanjor, T., Talirod, B., Tanjor, 
                                                                                                                                                       AC024224 165414 bp DNA linear PRI 29-MAY-2002 Homo sapiens 12 BAC RP11-133L14 (Roswell Park Cancer Institute Human BAC Library) complete sequence.
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Direct Submission
Submitted (26-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Submitted (28-FEB-2000) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA
                                                                                                                                                                                                                                                                                                                                                                Sukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
                                    661 AAGAATGACAGGGAGAAAGGAAAGAAGGGAAAATGTTGCCAAGGAAAAAAA 713
                                                                                                                                                                                                                                                                                                                                                                                        Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                     AC024224
AC024224.33 GI:21240476
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SSLEDSVTPTKAVKTTGKGIVKGRNLDSRGLILGAEAWGRGVKKNT"
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2 (bases 1 to 713)
Clark, H.F.
Direct Submission
Submitted (01-AUG-2003) Department of Bioinformatics, Genentech, Inc., I DNA Way, South San Francisco, CA 94080, USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               TITIGGGATITITAATITITCAAACACAGCAGAATGACATITITITITCTGTCACTATTATTAT
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92. .364
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/note="PR01159"
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product="CLECSF12"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match 100.
Best Local Similarity 100.
Matches 713; Conservative
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                                                                                                                                                          source
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complement(21141, 21950)
/rpt_family="LimC2"
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/rpt family="MLT1A1"
15398. 15421
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[. .1640]
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[21060]
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22220. .22101
                                                                                                                                                                                                                                                                                                                                  family="MER5B"
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ement(1700)
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complement(13288. .13579)
/rpt_family="MLT1A1"
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8877. .8904
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          6766. .6966
/rpt_family="Charlie2"
6980. .7102
                                                              /rpt_family="Charlie2"
7137. .7184
                                                                                                                  'rpt_family="AT_rich"
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complement(9745..100
                                                                                                                                                                                                                                                                                                                                                                                                  family="CT-rich"
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complement(15682..16
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11418. .11839
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family="(TA)n"
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1. .18101
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3.19811
family="LIM2"
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18102. .18318
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complement(20182
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complement(12307
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complement(13580
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  SEQUENCING READ COVERAGE:Sequencing is completed to a minimum standard of double strand coverage with a minimum of 2 clones and 2 reads with no ambiguities or 2 chemistries with a minimum of 2 clones and 3 reads with no ambiguities. If the sequence quality for a region does not meet this standard, it will be indicated in the annotation as Low Coverage.
                                                                                                                                                                                                               CLONE LENGTH: This sequence does not necessarily represent the entire insert of this clone. Overlapping regions of clones are only sequenced and submitted once, so the sequence for the remainder of the insert may be found in the record for the adjacent clones. Overlapping clones are noted at the beginning and end of the Features listing.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    note="Variable size of homopolymeric repeat in subclones"
                                                                                                                                                                                                                                                                                                                                                                                                                                 Indeptible transfer of the man and Mouse sequences.

Repeats are identified using RepeatMasker (A. Smit and P. Green, unpublished.) for Human and Mouse sequences.

Genes and Region of sequence similarity are identified by BLAST (Nuc. Acids Res., 25:3389-3402) similarity (expect < 1e-34) to the EST and cDNA sequences. Genes demonstrate at least two exons flanked by consensus splice sites that maintained sequence continuity across the splice junctions. Sequences that are not identical matches are annotated as similar.
                4 (bases 1 to 165414)
Worley,K.C.
Direct Submission
Submitted (29-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA
GO May 29, 2002 this sequence version replaced gi:21206012.
INFORMATION: http://www.hgsc.bcm.tmc.edu/ or email
                                                                                                                                                                                                                                                                                                                                                      ANNOTATION OF FEATURES:
STSs are identified using ePCR (Genome Res. 7:541-550) searches of a local database that includes entries from dbSTS, GDB, and local mapping efforts.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             QUALITY OF INDIVIDUAL BASES: This sequence meets stringent quality standards - estimated error rate less than 1 per 10,000 bases. Seports of lowest quality individual bases and measures of base quality are listed below. Description of the metrics can be found
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 http://gc.bcm.tmc.edu:8088/quality.info/genbank.annotation.html.
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TX 77030, USA
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/rpt_family="L1PA3"
complement (4558. .4629)
/rpt_family="MERSA"
4859. .4941
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 'organism="Homo sapiens"
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complement(6577. .6703)
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xref="taxon:9606"
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6420. .6491
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1 (bases 1 to 240864)

1 (bases 1 to 240864)

2 Muzny, D.M., Adame, C., Adio-Oduola, B., Ali-osman, F.R., Allen, C., Alsbrooks, S.L., Amaratunge, H.C., Are, J.R., Banks, T., Barbaria, J., Benton, J., Binage, K., Blankenburg, K., Bonnin, D., Bouk, J., Buhay, C., Benton, P., Burkett, C., Burrell, K.L., Byraht, N.P., Buhay, C., Burch, P., Burkett, C., Burrell, K.L., Byraht, N.C., Carron, T.F., Carter, M., Cavazos, S.R., Chacko, J., Chavazo, D., Chen, G., Chen, R., Chowdhry, I., Christopoulos, C., Cleveland, C.D., Coyle, M.D., Dathorne, S.R., David, R.N., Davia, M.L., Davis, C., Davy-Carroll, L., Dederich, D., Dablador, D., Bandor, D., Bolandor, D., Bandor, D., Bandor, D., Bandor, D., Bandor, D., Bandor, D., Flagg, N., Ford, J., Garza, N., Gall, R., Gorrell, J.H., Guevara, W., Gunarathe, P., Hale, S., Hamilton, K., Harris, C., Harris, K., Hart, M., Havlak, P., Hawes, A., Hernandez, J., Hernandez, J., Howara, S., Hober, J., Holloway, C., Hollins, B., Homsi, F., Howard, S., Huber, J., Hulyk, S., Hume, J., Jackson, L.E., Jacobson, B., Jia, Y., Johnson, R., Jolivet, S.,
                                                116999 GTATCCCTGTCATATATACAATAAGGTGAAATTATAAGTACCCTATGCAGTTGGCTGGAC 116940
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Homo sapiens clone RP11-13C13, RP11-656E20, *** SEQUENCING IN
ACOGSSIO ACOGSSI 4.7. 16 unordered pieces.
ACCOGSIO ACOGSSI 4.7. 1700 ACCOGSIO ACCOCSIO ACCOGSIO ACCOCSIO ACCOC
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GITGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCTA 180
                                                                                                                                                                        CTTTCTATCAAGAAATAAAGAGAACCACAGTCAACCCACACAATCATCTTTAGAAGACAG
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Homo sapiens (human)
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Joudah,S., Karlsson,E., Kelly,S., Khan,U., King,L., Korvah,J.,

Kovar,C., Kratovic,J., Kureshi,A., Landry,N., Leal,B. Lewis,L.C.,

Lewis,L., Li,J., Li,Z., Lichtarge,O., Lieu,C., Liu,J., Luna,R.,

Ma,J., Maheshwari,M., Mapua,P., Martin,R., Martindale,A.,

Matrinez,B., Massey,E., Mawhiney,E., McCheod,M.P., Meador,M.,

Matrinez,B., Massey,E., Mawhiney,E., Mitchell,T., Mohabbat,K.,

Morgan,M., Morris,S., Moser,M., Neal,D., Newtson,J., Newtson,N.,

Nguyen,A., Nguyen,N., Nguyen,N., Orkenson,E., Nwokenkwo,S.,

Oguh,M., Okwuonu,G., Oragunye,N., Ortedo,R., Pace,A., Payton,B.,

Petery,J., Peters,L., Pickens,R., Primms,E., Pu,L.L.,

Quiles,M., Ren,Y., Rives,M., Rojas,A., Shen,H., Shocshtari,N.,

Ruiz,S., Savery,G., Scherer,S., Scott,G., Shen,H., Shocshtari,N.,

Sisson,I., Sodergren,E., Sonake,T., Sparks,A., Stanley,H.,

Tang,H., Taneey,J., Taylor,C., Taylor,T., Talfrod,B., Thomas,N.,

Thomas,S., Usmani,K., Vasquez,L., Vera,V., Villalon,D., Vinson,R.,

Matlington,S., Williams,G., Walten,R., Washington,C.,

Morley,K., Wu,C., Wu,Y., Wu,Y.F., Zhou,J., Zorrilla,S., Nelson,D.,

and Gibbs,R.,

M. Dublished

L. Unpublished
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Chemistry: Dye-terminator Big Dye: 23% of reads
Assembly program: Phrap; version 0.990329
Consensus quality: 71594 bases at least Q30
Consensus quality: 93014 bases at least Q30
Consensus quality: 110702 bases at least Q20
Estimated insert size: 229041; sum-of-contigs estimation
Bstimated insert size: 331000; agarose-fp estimation
Quality coverage: 0.6x in Q20 bases; agarose-fp estimation
Quality coverage: 0.9x in Q20 bases; sum-of-contigs estimation
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Submitted (05-FEB-1999) Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA ON 58p 14, 2000 this sequence version replaced gi:10086352.
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gap of unknown length
contig of 23597 bp in length
gap of unknown length
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Center project name: J-33
Center clone name: RP11-13C13, RP11-656E20
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Web site: http://www.hgsc.bcm.tmc.edu/
Contact: hgsc-help@bcm.tmc.edu
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Worley, K.C.
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163754:
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202627:
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194802:
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us-09-989-293a-376.rge

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211151 TCTCGCCCTTACACGAATTGACATTTTAAATGCGATACAGTTAGAATAGGAAATATGACA 211092
                                                                                                                                                                                                                                  AC138620 221471 bp DNA linear HTG 25-FEB-2003 Mus musculus chromosome UNK clone RP23-250E1, WORKING DRAFT SEQUENCE, 13 unordered pieces.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Submitted (12-JAN-2003) Genome Sequencing Center, 4444 Forest Park Parkway, St. Louis, MO 63108, USA 3 (bases 1 to 221471) McPherson, J.D. and Waterston, R.H.
                                                                                                  Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
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                                                                   TTAGAAAGGAAGAATGACAGGAGAAAGGAAAGGAAAATGTTGCCAAGGAAAA
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Parkway, St. Louis, MO 63108, USA
On Peb 25, 2003 this sequence version replaced gi:27657609.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequencing vector: plasmid; 10%
Sequencing vector: plasmid; 100%
Chemistry: Dye-primer ET; 0% of reads
Chemistry: Dye-primer ET; 0% of reads
Assembly program: Phrap; version 0.990319
Consensus quality: 217739 bases at least Q40
Consensus quality: 218078 bases at least Q20
Consensus quality: 218078 bases at least Q20
Insert size: 193000; agarose-fp
Insert size: 1931013; sun-of-conitiss
Quality coverage: 14.88 in Q20 bases; sum-of-contigs
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Center code: WUGSC
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HTG; HTGS PHASE1; HTGS_EVLLTOP
Mus musculus (house mouse)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       1 (bases 1 to 221471)
Mcherson, J.D. and Waterston, R.H.
The sequence of Mus musculus clone
Unpublished
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McPherson, J.D. and Waterston, R.H.
Direct Submission
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3424
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AC138620/c
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ACTITC-TATCAAGAAATAAAGAGAACCACAGT-CAACCCACACAATCATCTTTAGAAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  211571 ACTITICTIATICAAGAATAAAGAGAACCACAGTCCACCCCACACATTCTTTAGAAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   CAGTGTGACTCCTACC-AAAGCTGTCAAAACCA-CAGGCAAGGGCATAGTTAA-AGGACG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   211451 GARICTIGACTCAAAGAGGGTTAATTCTTGGTGCTGAAGCCTGGGGCAGGGGTGTAAAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    354 AAAACACTTAGATTCAATGATTGTAAATTTAAGGCAAATACACATATTAGTATTACCTTA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    GTGTAATGTATCCCTGTCATATATACAATAAGGTGAAATTATAAGTACCCTATGCAGTTG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      GCTGGACAGTTCTAAATTGGACTTTATTAATTTTTAAAATCAGTAACTGATTTATCACTG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        GCTATGTGCTTAGATCTACAGGAGATCATATAATTTGATACAAATAAAAGAAAAGTGTTC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   211631 TGTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TGTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGGCT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 240864;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    6
8: contig of 60.2.
58: gap of unknown length
85: contig of 13027 bp in length
885: gap of unknown length
1342: contig of 8257 bp in length
1342: contig of 8257 bp in length
13272: gap of unknown length
13725: gap of unknown length
137459: contig of 1334 bp in length
234459: contig of 1734 bp in length
23459: gap of unknown length
235921: gap of unknown length
235921: gap of unknown length
235921: gap of unknown length
235941: contig of 1225 bp in lengt
23746: gap of unknown length
238417: contig of 1171 bp in lengt
235949: contig of 1132 bp in length
23649: contig of 1132 bp in length
23649: contig of 1132 bp in length
23649: contig of 1115 bp in length
2504964: contig of 1115 bp in length
2504064: contig of 1115 bp in length
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              82.8%; Score 590.2; DB 2; 96.2%; Pred. No. 6.8e-111; iive 0; Mismatches 18;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           /clone="RP11-13C13, RP11-656E20"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                /organism="Homo sapiens"
/mol_type="genomic DNA"
/db_xref="taxon:9606"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Matches 690; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Local Similarity
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Allen, C., Allen, H., Alsbrooks, S., Amin, A., Anguiano, D., Anyalebechi, V., Aoyagi, A., Ayodeji, M., Baca, E., Baden, H., Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F., Bayanlo, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M., Barater, N., Carter, K., Cavazos, I., Ceasar, H., Center, A., Chacko, J., Chaves, D., Chen, G., Chen, Y., Chen, Z., Chen, Z., Chacko, J., Chaves, D., Chen, G., Chen, Y., Chen, Z., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D., Delgado, O., Denson, S., Deramo, C., Dinh, Y., Dinh, H., Divya, K., Draper, H., Duyan-Rocha, S., Durn, A., Durbin, K., Duval, B., Eaves, K., Ernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Gabrai, A., Garcia, R., Garner, T., Garza, M., Gebregeorgis, E., Geer, K., Gill, R., Gracia, A., Garner, T., Garza, M., Gunaratne, P., Haaland, M., Hamel, C., Hamilton, C., Hamilton, K., Harnandez, M., Havlak, P., Hawes, A., Henderson, N., Hernandez, J., Jackson, A., Hollins, B., Howells, S., Hullyk, S., Hullyk, S., Kally, S., Liu, J., Li, Z., Liu, J., Kovar, C., Kovar, C., Kovar, C., Kowis, C., Liu, J., Lewis, C., Liu, J., Lewis, L., Li, Z., Liu, J.,
                ------Trircaciracraaaagaacagraacagcrcrirarcrc 86612
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           86494 GGGAAAGGAAATATAAGGGGAGAAATGAGGAAGGGACACTGAAGACTGTAAATGCTAATAA 86435
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AC112033 273867 bp DNA linear HTG 10-OCT-2002
Rattus norvegicus clone CH230-94G2, *** SEQUENCING IN PROGRESS ***,
2 unordered pieces.
                                                                                                                CGGAATCTTGACTCAAGAGGGTTAATTCTTGGTGCTGAAGCCTGGGGGCAGGGGTGTAAAG 352
                                                                                                                                                                                                                                                                                                                                                                       AGTGTAATGTATCCCTGTCATATATACAATAAGGTGAAATTATAAGTACCCTATGCAGTT 472
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Rodentia, Sciurognathi, Muridae, Murinae,
                                                                                                                                                                                                                                                                                                         86763 -AATATGTTTACATCTACTAACAGTGAATTCAAGGCAAGCAGACCTAACAGAATTCCCTT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            86704 AGCATAACATA-CCCTGCCAGACAGCTGGTAAAGTGGCATCACAAATCCACAGCTC----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              473 GGCTGGACAGTTCTAAATTGGACTTTATTAATTTTAAAATCAGTAACTGAT-TTATCAC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         TCTCTCCCCTTACAGAATTGACATTTTAAATGCGATACAGTTAGAAATAGGAAATATGACA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 86551 TCAGTGTCCTTATGGAACTGATATTTTACAGTGAT---GTTAGAATAAGATAGATGGA
                                                                                                                                                                                                                                             AAAAACACTTAGATTCAATGATTGTAAATTTAAGGCAAATACACATATTAGTATTACCTT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           TGGCTATGTGCTTAGATCTACAGGAGATCATATAATTTGATACAAATAAAAGAAAAGTGT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                652 TTAGAAAGGAAGAATGACAGGGAGAAAGGAAAGGAAAAGGGAAAATGTTGCCAAGGAAAAAA
                                                                                                                                                                     86815 GGGATCCCGGATTCCAGAGGTTCATTTCCTGGTGCTGAGGCCTGGGGTGGTGGG
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HTG; HTGS PRASE1; HTGS DRAFT; I
Rattus norvegicus (Norway rat)
Rattus norvegicus
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KEYWORDS
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ORGANISM
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AC112033
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 86992 GTTCGTGACGTATGAAGCATTTTGGTGACACTATTCAGGGAGAAATCCAGAGGAGAAAGA 86933
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           86932 CAACTICCTATCAAGAAATAAAGAGAACCACA---AGCCCACAGAATCATCTTTAGATGA 86876
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             86993
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     118 ATTGTTGGTATGTGAAGCTATTTGGAGATCCAATTCAGGAAGCAACACATTGGAGAATGG 177
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 1 AATATATCATCTATTTATCATTAATCAATAATGTATTCT-TTTATTCCAATAACATTTGG
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                87052 GIGCIGGCAITICCAITITIAAACACAGAAIGACITICITITITICIGICACIATIGCI
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CTACTITCTATCAAGAAATAAAGAGAACCACAGTCAACCCACACAATCATCTTTAGAAGA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         238 CAGTGTGACTCCTACCAAAGCTGTCAAAACCACAGGGCAAGGGCATAGTTAAAGGA----
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
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1091 20220: contig of 6130 bp in length 20221 33395: contig of 13075 bp in length 33495: gap of unknown length 3495: gap of unknown length 4496 46731: contig of 13275 bp in length 4596 46731: app of unknown length 6732 46831: gap of unknown length 62519: contig of 12587 bp in length 62519: contig of 12587 bp in length 80892: contig of 18274 bp in length 80892: contig of 18274 bp in length 80892: contig of 22320 bp in length 103312: contig of 22320 bp in length 125914: contig of 22320 bp in length 125914: contig of 22320 bp in length 11559: contig of 45445 bp in length 11559: contig of 49912 bp in length 11559: contig of 49912 bp in length 11559: contig of 49912 bp in length 11550: contig of 49912 bp in le
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Pred. No. 2.1e-28;
0; Mismatches 215; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        .610 .3423
/note="assembly_name:Contig21"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             note="assembly_name:Contig23"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         .0321. .33395
'note="assembly_name:Contig25"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        3496. .46731
note="assembly_name:Contig26"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                16832, .62518
/note="assembly_name:Contig27"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               62619. .80892
/note="assembly_name:Contig28"
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26015. 171459
hote="assembly_name:Contig31"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          note="assembly_name:Contig19"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              note="assembly_name:Contig22"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               note="assembly_name:Contig24"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              note="assembly_name:Contig29"
103413. .125914
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     organism="Mus musculus"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            'mol_type="genomic DNA"
'db_xref="taxon:10090"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               'clone="RP23-250E1"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              chromosome="UNK"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       0993. .103312
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Best Local Similarity 63.2%;
Matches 456; Conservative (
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(Dases 1 to 273867)

Rat Genome Sequencing Consortium.

Direct Submission

Olivect Submission

All Submission

All Submission

All Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor plaza, Houston, TX 77030, USA

On Oct 10, 2002 this sequence version replaced gi:21737395.

The sequence in this assembly is a combination of BAC based reads and whole genome shotgun sequencing reads assembled using Atlas in the feature table below represents a scaffold in the Atlas assembly (a 'contig-scaffold') Within each contig-scaffold, individual sequence contigs are ordered and oriented, and separated by sized gaps filled with Ng to the estimated size. The sequence may extend beyond the ends of the clone and there may be sequence contigs within a contig-scaffold that consist entirely of whole genome shotgun sequence reads. Both end sequences and whole genome shotgun sequence reads. Both end sequences and whole genome table.
Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J.,
Lourensuhewa, L., Loulseged, H., Lozado, R.J., Lu, X., Ma, J.,
Mahesbwari, M., Mahindartne, M., Mahmoud, M., Malloy, K., Mangum, A.,
Mangum, B., Mayua, P., Martin, K., Martinez, E.,
Mangum, B., McLeod, M.P., McNeill, T.Z., Meenen, B.,
Mangum, B., McLeod, M.P., McNeill, T.Z., Meenen, B.,
Mangum, M., Morris, K., Morris, S., Murphy, M., Nair, L.,
Nankervis, C., Neal, D., Newton, N., Nguyen, N., Norris, S.,
Nwaokelemeh, O., Okwuon, G., Olarnpunsagoon, A., Pal, S., Parks, K.,
Pasternak, S., Paul, H., Perez, J., Prannkoch, C.,
Plopper, P., Poinderer, A., Popovic, D., Primus, E., Pu, L.-L.,
Plopper, P., Poinderer, A., Popovic, D., Primus, E., Pu, L.-L.,
Pasternak, S., Paul, H., Ren, Y., Reuter, M., Regier, M., Reigh, R.,
Rives, C., Rodkey, T., Rachlin, E., Secher, S., Shatsman, S., Shen, H.,
Sanders, W., Savery, G., Scherer, S., Sotelle, R., Shatsman, S., Shen, H.,
Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smajs, D.,
Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smajs, D.,
Shedy, T., Thomas, N., Thomas, N., Trejos, Z., Usmani, K.,
Valas, R., Vera, V., Villasana, D., Waldron, L., Walker, B., Wang, J.,
William, G., Willson, M., Warren, R., Wei, Y., Walker, B., Wang, J.,
Williams, G., Willson, M., Warren, R., Wei, Y., Yoon, L., Yoon, V.,
Williams, G., Willson, M., Walker, W., Ven, J., Yoon, L., Yoon, V.,
Williams, G., Willson, M., Walker, S., Dunn, D., von,
Niederhausern, A., Weiss, R., Smith, D. R., Holt, R., Smith, H.O.,
Weinstock, G. and Gibbs, R.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   NOTE: Estimated insert size may differ from sequence length (see http://www.ngsc.bom.tmc.edu/docs/Genbank draft_data.html) NOTE: This sequence may represent more than one clone. NOTE: This is a 'working draft' sequence. It currently
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Submitted (19-FEB-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Quality coverage: 6x in Q20 bases; sum-of-contigs estimation
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Consensus quality: 244419 bases at least Q40
Consensus quality: 247444 bases at least Q30
Consensus quality: 249002 bases at least Q20
Estimated insert size: 254595; sum-of-contigs estimation
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Center code: BCM
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Center project name: GPBG
Center clone name: CH230-94G2
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Direct Submission
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AUTHORS TITLE JOURNAL

COMMENT

REFERENCE

REFERENCE AUTHORS TITLE JOURNAL

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46206 AGGIGGCICCCICCAAGCAICCCAAACIACGAGGCAAGGCAAGAACIAAAGGAGCAAIGG 46265
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Homo sapiens beta-glucan receptor isoform H (BGR) mRNA, complete cofs, alternatively spliced.
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consists of 2 contigs. The true order of the pieces is not known and their order in this sequence record is arbitrary. Gaps between the contigs are represented as runs of N, but the exact sizes of the gaps are unknown. This record will be updated with the finished sequence as soon as it is available and the accession number will be preserved.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 20.7%; Score 147.6; DB 2; Length 273867; 71.7%; Pred. No. 1.4e-20; ive 0; Mismatches 89; Indels 10; 0
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                                                                                                                                                                                     1 271836: contig of 271836 bp in length 271837 271936: gap of unknown length 271937 273867: contig of 1931 bp in length. Location/Qualifiers 1. 273867 contig of 1931 bp in length. J. 273867 contig of 1931 bp in length. Jorganism="Rattus norvegicus"
                                                                                                                                                                                                                                                                                                                                                                                                                                                     /note="wgs_end_extension
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7780. 8635
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clone_end:T7
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/clone="CH230-94G2"
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7780. .8493
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                        /product="beta-glucan receptor isoform H"
/protein id="AAL11718.1"
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LIAVILGILCLVILVIAVVLGTMGTGQFLKDLSFLNNRRKLFGDPIQEATHWRMATFY
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                            Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi, Mammalia; Butheria, Primates, Catarrhini, Hominidae, Homo.
1 (bases 1 to 881)
Willment, J. A. Gordon, S. and Brown, G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
J. Biol. Chem. 276 (47), 43818-43823 (2001)
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Bukaryotas, Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Butheria; Primates; Catarrhini; Hominidae; Homo.

1 (bases 1 to 625)

Willment, J.A., Gordon, S. and Brown, G.D.

Characterization of the human beta -glucan receptor and its
alternatively; spliced isoforms
21570237

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                                                                                                                                                                          Dises 1 to 881)
Willment, J.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JUD-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
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/note="C-type lectin BGRH; alternatively spliced"
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/chromosome="12"
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L. .881
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Homo sapiens (human)
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Best Local Similarity
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/trānslation="MEYHPDLENLDEDGYTQLHFDSQSNTRIAVVSEKGSCAASPPWR
LATVIGGICCLVILVIAVYLGTWAITRSNSGSNTLENGYFLSRNKENHSQPTQSSLED
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SNELISDQNHSYPRKPISKICMDSRVSHL"
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; alternatively spliced"
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2 (bases 1 to 625)
Willmant,J.A., Gordon,S. and Brown,G.D.
Direct Submission
Direct Submission
Size (18-J1-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
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                                                                                                                                                                                                                                                                                                      tissue_type="peripheral blood leukocyte"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       product="beta-glucan receptor isoform C"
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unclassified.
(Loases 1 to 741)
Shuji, H., Shoji, F., Kazunori, N. and Yasushi, A. Novel membrane protein and DNA thereof
Exent: JP 1999001497-A 2 06-JAN-1999;
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                                                                                                                                                                          /organism="Homo sapiens"
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JP 1999001497-A/2
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/protein id="AAL:1711.1"
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1. (bases 1 to 744)

Willment, J. A., Gordon, S. and Brown, G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
J. Biol. Chem. 276 (47), 43818-43823 (2001)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AF400595 744 bp mRNA linear PRI 19-NOV-20
Homo sapiens beta-glucan receptor isoform A (BGR) mRNA, complete
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Willment, J.A., Gordon, S. and Brown, G.D.
Direct Submission
Submitted (18-JUL-2201) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK
Location/Qualifiers
A61K37/02, A61K37/02, C12N15/00, (C12N15/00, C12R1:91) CC
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                                                                                                                                                                                                           Length 741;
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1; alternatively spliced"
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                                                                              organism='Unidentified'
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al Similarity 100.0%; Pred. No. 1.2e-18;
140; Conservative 0; Mismatches 0;
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                                                                                          location/Qualifiers
1. 741
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/chromosome="12"
/map="12p13"
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 PC A61K37/02,A61K37/0
Strandedness: Double;
CC Topology: Linear;
FH Key Lo
FT source 1.0
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/product=1lectin-like receptor 1"
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SNBLGFIVKQVSSQPDNSFWIGLSRPQTEVWLWEDGSTFSSNLFQIRTTATQENPSP
NCVWIHVSVIYDQLCSVPSYSICEKKFSM"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Hermanz-Falcon, F., Arce, I., Roda-Navarro, P. and Fernandez-Ruiz, E. Cloning of human DECTIN-1, a novel C-type lectin-like receptor gene expressed on dendritic cells
Immunogenetics 53 (4), 288-295 (2001)
LIAVILGILCLVILVIAVVLGTMAIMRSNSGSNTLENGYFLSRNKENHSQPTQSSLED
SVEPKAVKTTGVLGSPCPFWAITENSKSCYLESMSLNSWDGSKRQCWQLGSNILKIDS
SNELGFIVKQVSSQPDNSFWIGLSRPQTEVWLWEDGSTFSSNLFQIRTTATQENPSP
NCVWHHYSVIYDQLCSVPSYSICEKKFSM"
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 744)
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Homo sapiens lectin-like receptor 1 (DECTIN1) mRNA, complete cds.
AY026769
                                                                                                                                                                                                                                                                                                                                        202 GCTATTTGGAGATCCAATTCAGGAAGCAACATTGGAGAATGGCTACTTTCTATCAAGA
                                                                                                                                                                                                                                                                                                                                                                                                  194 AATAAAGAGAACCACAGTCAACCCACACAATCATCTTTAGAAGACAGTGTGACTCCTACC
                                                                                                                                                                                                                                                                                   134 GCTATTTGGAGATCCAATTCAGGAAGCAACACTTGGAGAATGGCTACTTTCTATCAAGA
                                                                                                                                                                                                                              Gaps
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Direct Submission

Submitted (01-FBB-2001) Biologia Molecular, Hospital de la Princesa, Diego de Leon 62, Madrid 28006, Spain

Herranz-Falcon, P., Arce, I. and Fernandez-Ruiz, E.

Direct Submission

Submitted (05-OCT-2001) Biologia Molecular, Hospital de la Princesa, Diego de Leon 62, Madrid 28006, Spain

Sequence update by submitter

Sequence update by submitter

On Oct 5, 2001 this sequence version replaced gi:14278818.
                                                                                                                                                                                                                              0;
                                                                                                                                                                       Length 744;
                                                                                                                                                                                                                              Indels
                                                                                                                                                                         19.6%; Score 140; DB 9; I
100.0%; Pred. No. 1.2e-18;
cive 0; Mismatches 0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          sapiens (human)
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                                                                                                                                                                                                                                 Conservative
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ORIGIN

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SVTPTKAVKTTGVLSSPCPPRWIIYEKSCYLESMSLNSWDGSKRQCWQLGSNLIKIDS
SNELSLTLLPKLECSEAATSQAQVILPPQLPB"
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi, Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
I (basea 1 to 787)
Willment, J. A., Gordon, S. and Brown, G.D.
Characterization of the human beta -glucan receptor and its alternatively spliced isoforms
J. Biol. Chem. 276 (47), 43818-43823 (2001)
                                                                              134 GCTATTTGGAGATCCAATTCAGGAAGCAACACTTTGGAGAATGGCTACTTTCTATCAAGA
                                                                                                                                                           AATAAAGAGAACCACAGTCAACCCACACCAATCATCTTTAGAAGACAGTGTGACTTCCTACC
                                          Gaps
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Willment, J.A., Gordon, S. and Brown, G.D.

Direct Submission

Submitted (18-JUL-2001) Sir William Dunn School of Pathology,
Oxford University, South Parks Road, Oxford OX1 3RE, UK

Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   /note="C-type lectin BGRG; alternatively spliced"
                                            0;
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      Length 744;
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                                          0; Indels
    19.6%; Score 140; DB 9; I
100.0%; Pred. No. 1.2e-18;
iive 0; Mismatches 0;
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/organism="Homo sapiens"
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/chromosome="12"
/map="12p13"
1. .787
/gene="BGR"
                                                                                                                                                                                                                                                            322 AAAGCIGICAAAACCACAGG 341
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  Query Match
Best Local Similarity 100.
Matches 140; Conservative
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DEFINITION
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KEYWORDS
SOURCE
ORGANISM
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AUTHORS
TITLE
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AF400601
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AUTHORS
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PUBMED
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        QY
        194 AATAAAGAACCACACACCACACAATCATCTTTAGAAGACAGTGTGACTCCTACC 253

        Db
        262 AATAAAGAACCACAGCAACCATCATCTTTAGAAGACAGTGTGACTCTTACC 321

        QY
        254 AAAGCTGTCAAACCACAGG 273

        Db
        322 AAAACTGTCAAAACCACAGG 341
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Search completed: December 2, 2004, 04:01:55 Job time: 4007.48 secs

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473.
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Human cDNA encoding a secreted/transmembrane protein, SEQ ID
US2003032155-A1.
               100.0%; Score 713; DB 8; Length 713; 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
10.2003022187-A1.
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Novel human secreted or transmembrane protein PRO1124 DNA, US20030227985-A1.
06-FEB-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ACD42031 standard; cDNA; 713 BP.
Human secreted/transmembrane protein (PRO) cDNA #237.
US2003036179-A1.
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(GETH ) GENENTECH INC.
lery Match
lery Match
100.0%; Score 713; DB 8; I
                                                                                                                                                                                                         Score 713; DB 8;
Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Human sectreted/transmembrane protein cDNA, #154.
US2002142951-A1.
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Pred. No. 3.3e-155;
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No. 3.3e-155
                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABX80868 standard; cDNA; 713 BP.
Human secreted/transmembrane protein cDNA, #154.
US2003027162-A1.
                                                                                                       cDNA encoding human PRO polypeptide #237 US2003036180-A1.
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cDNA encoding human PRO1159 polypeptide.
US2002127576-A1.
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Pred.
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20-FEB-2003.
(GETH ) GENENTECH INC.
100.0%; SC
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13-FEB-2003.
(GETH ) GENENTECH INC.
100.0%; Sr
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
            Query Match
Best Local Similarity
RESULT 6
                                                                                                                                                                                                                                                                             ABX89377 standard;
DNA encoding novel
US2003017563-A1.
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Best Local Similarity
RESULT 7
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                                                                                ACA03839 standard;
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GenCore version 5.1.6

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scember 2, 2004, 00:55:08 ; Search time 534.388 Seconds
(without alignments)
7003.964 Million cell updates/sec
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SUMMARIES
                                                                                                                                                                   Sequence:
Scoring table:
BENTITY NUC
Gapor 10.6, Gapext 1.0
Searched:
Total number of hits satisfying chosen parameters:
Minimum DB seq length:
Maximum Match 0%
Post-processing:
Minimum Match 100%
Maximum Match 100%
Database:
N Geneseq.1990s:*
Sequence:
Maximum Match 100%
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US200300321-A1.
02-JAN-2003.
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Human PRO1159 (UNQ589) nucleotide sequence SEQ ID NO:376.
WC200454-A1.
07-DEC-2000.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human PRO1159 protein UNQ589 encoding cDNA SEQ ID NO:272. WO200053758-A2.
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e encoding for PRO1159 polypeptide
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Membrane-bound protein PRO1159 encoding cDNA.
W09963088-A2.
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geneseqn2003ds:*
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07-JUN-2001.
(GETH ) GENENTECH INC.
100.0%;
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713
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 3
ID ACA64409 standard; ci
DE Novel human secreted
PN US200300351-A1.
PD 02-JAN-2009.
PA (GETH ) GENENTECH INC
                                                    December
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Human cDNA sequence
WO200140466-A2.
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Best Local Similarity
RESULT 2
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Perfect score:
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(GETH) GENENTECH INC.

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PD 30-JAN-2003.
PA (GETH) GENENTECH INC.
Query Match 100.0%; Score 713; DB 9; Length 713;
Best Local Similarity 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003068796-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003017476-A1.
23-JAN-2003.
                                               Length 713;
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Human secreted and transmembrane polypeptide PRO1159 cDNA
22002197615-A1.
                                                                                                                                                                                                                   ACA68006 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PR01159
US2002177164-A1.
28-NOV-2002.
(GETH ) GENENTECH INC.
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PA (GETH ) GENENTECH INC.

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Best Local Similarity 100.0%; Pred. No. 3.3e-155;

RESULT 22
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
                                               100.0%; Score 713; DB 8;
100.0%; Pred. No. 3.3e-155;
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US2003073212-A1.
                                                                                            ABX17151 standard; cDNA; 713 BP. Whman FW Dolynucleotide #118. US2002123463-A1. 05-SEP-2002.
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Human PRO polynucleotide #237.
US2003054517-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity .RESULT 20
ID AD445992 standar
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Best Local Similarity
RESULT 19
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ADA11573 standard; cDNA; 713 BP.
Human cDNA encoding secreted/transmembrane polypeptide PRO1159.
US2003054404-A1.
20-MAR-2003.
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(GETH) GENENTECH INC.
ery Match 100.0%; Score 713; DB 9; Length 713;
ery Match 100.0%; Pred. No. 3.3e-155;
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Human cDNA encoding secreted/transmembrane protein, PRO1159.
US2003059831-A1.
27-MAR-2003.
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Muman CDNA encoding secreted/transmembrane protein PRO1159.
US2003008297-A1.
09-JAN-2003.
(GERH) GENENTECH INC.
100.0%; Score 713; DB 9; Length 713
ety Match
st Local Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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(GETH) GENENTECH INC.
(ery Match 100.0%; Score 713; DB 9; )
(ery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
100.0%; Score 713; DB 9; 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
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                                                                   cDNA encoding human PRO polypeptide #237 US2003082704-A1.
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US2003073215-A1.
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17.APR-2003.
(GTH ) GENENTECH INC.
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(GETH) GENENTECH INC.
100.0%;
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RESULT 32
Query Match
Best Local Similarity
RESULT 25
ID ADB28022 standard;
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Best Local Similarity
RESULT 30
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Best Local Similarity
RESULT 27
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Human cDNA encoding secreted/transmembrane protein PRO1159 US2003059832-A1.
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Novel human secreted and transmembrane protein PR01159
US2003082691-A1.
01-MAY-2003
(GETH ) GENENTECH INC.
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                                                                100.0%; Score 713; DB 9; 100.0%; Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
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Best Local Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
100.0%; Score 713; DB 9; 100.0%; Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
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Human PRO polynucleotide SEQ ID NO 473.
US2003077713-A1.
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Human PRO polynucleotide #237.
US2003077722-A1.
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Human PRO polynucleotide #237.
US2003082701-A1.
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US2003068798-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                           17-APR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 49
                                                                                                                                                                                                                                 Local Similarity
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Best Local Similarity
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Best Local Similarity
RESULT 50
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           US2003073211-A1.
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RESULT 44
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(GETH ) GENENTECH INC.
ery Match 100.0%; Score 713; DB 9; Length 713;
ery Match 100.0%; Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
lery Match 100.0%; Score 713; DB 9; Length 713;
RESULT 34
ID ADA85949 standard; CDNA; 713 BP.
DB Novel human secreted and transmembrane protein PRO1159 CDNA
PN US2003082693-A1.
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(GETH ) GENENTECH INC.
ery Match
100.0%; Score 713; DB 9; Length 713;
ery Match
100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA,
US2003087345-A1.
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                                                                                                                     Length 713
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082694-A1.
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Muman cDNA encoding secreted/transmembrane protein PRO1159 US2003054359-A1.
                                                                    GETH ) GENENTECH INC.

(GETH ) GENENTECH INC.

(ery Match

100.0%; Score 713; DB 9; I
                                                                                                                                                                                                                                                 100.0%; Score 713; DB 9; 100.0%; Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.

(GETY Match 100.0%; Score 713; DB 9;

(ery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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08-MAY-2003.

(GETH ) GENENTECH INC.

100.0%; Score 713; DB 9;

lery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
                                                                                                                                                                    ADA17904 standard; cDNA; 713 BP. cDNA encoding human PRO1159 polypeptide. US2003054987-A1.
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                                                                                                                                                                                                                                                                                                  AAA97161 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003082705-A1.
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A (GETH ) GENENTECH INC.
Query Match
Best Local Similarity 100.0%;
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Best Local Similarity
RESUL 42
ID ADB14961 standard; cl
Pp Human PRO polynucleoi
PN US2003087351-A1.
PD 08-MAY-2003.
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Best Local Similarity
RESULT 39
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Best Local Similarity
RESULT 36
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Best Local Similarity
RESULT 35
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Best Local Similarity
RESULT 38
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Best Local Similarity
RESULT 41
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RESULT 40

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#DA93568 standard; cDNA; 713 BP.

Human PRO polynucleotide #237.

US2003077721-A1.

(GETH ) GENENTECH INC.

ery Match

100.0%; Score 713; DB 9; Length 713;
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Pred. No. 3.3e-155;
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                    Length 713;
                                                                ADA85397 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082695-A1.
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(GETH) GENENTECH INC.
ery Match 100.0%; Score 713; DB 9; Length 713;
ery Match 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082708-A1.
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Whwan cDNA encoding secreted/transmembrane protein PRO1159
US2003059780-A1.
27-MAR-2003.
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24-APR-2003.
(GETH ) GENENTECH INC.
100.0%; SCORE 713; DB 9; L
LERY MATCh 100.0%; Pred. No. 3.3e-155;
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(GETH) GENENTECH INC.

(GETH) MAtch 100.0%; Score 713; DB 9; L

ery Match 100.0%; Pred. No. 3.3e-155;
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PA (GETH ) GENENTECH INC.
Query Match
Best Local Similarity 100.0%; Pred. No. 3.3e-155;
RESULT 53
                                                                                                                                                                                                                                                                                                    cDNA encoding human PRO polypeptide #237
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              BP.
ID NO 473.
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Human PRO polynucleotide #237.
US2003073210-A1.
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                                                                                                                                                                                                                                                                                              ADB30101 standard; cDNA; 713 BP.
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Human PRO polynucleotide #237.
US2003082761-A1.
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Human PRO polynucleotide SEQ
US2003077715-A1.
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(GETH ) GENENTECH INC.
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.59780-Al.
.4ry Match
Best Local Similarity in RESULT 59 6 stande.
DD ADA47096 stande.
PN US2003"
PD 17-
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Judy,

Lery Match
Best Local Similarity
RESULT 58
ID ADA38817 stand
DE Human CDW
PN US200
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                                                                                                                                      Query Match
Best Local Similarity
RESULT 54
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Best Local Similarity
RESULT 56
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Best Local Similarity
RESULT 60
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Length 713;
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                                                                                                                                                                                                                                                                                                                                                                       Length 713;
                                                                                                                                                                                                                                                                                                                   Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003060407-A1.
27-MAR-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                         ADA61135 ...
Homo sapiens.
Homo sapiens.
13 MAR-2003.
(GETH ) GENENTECH INC.
100.0%; Score 713; DB 9; Lv.
100.0%; Pred. No. 3.3e-155;
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PA (GETH ) GENENTECH INC.
Query Match
Best Local Similarity 100.0%; Pred. No. 3.3e-155;
RESULT 70
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100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                        ADB31205 standard; cDNA; 713 BP.
CDNA encoding human PRO polypeptide #237.
US200306386-Al.
22-MAY-2003.
(GETH) GENENTECH INC.
ery Match
st. Local Similarity 100.0%; Score 713; DB 9; I
                                                                                                                                                                                                                                                                                                                                                                         100.0%; Score 713; DB 9; 100.0%; Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
Best Local Similarity 100.0%; Pred. No. 3.3e-155; RESULT 62
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADB24280 standard; CDNA; 713 BP.
Human PRO polynucleotide SEQ ID NO 473.
US2003077714-A1.
                                       _ADB26918 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide US2003092147-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human PRO polynucleotide #237.
US2003082759-A1.
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                                                                                                                                                                                                                                                                                                           ADA92938 standard; cDNA; 713 BP.
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RESULT 71
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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LUD3.

LUCAL MATCH

BEST LOCAL Similarity

RESULT 64

ID ADA92938 stand

DE Human Chw.

PN US2nd

PD
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Best Local Similarity
RESULT 69
                                                                                                                                                                                                                                                                                                                                                                                          Best_Local Similarity
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BP.
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100.0%;
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                          27-MAR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
                                                                          Local Similarity
                                                                                                              ADA22499 standard;
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ADB27470 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Local Similarity
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ID ADB22403 standard;
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ID ADA67094 standard;
            US2003059909-A1.
                                                                                                                                                                27-FEB-2003.
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(GETH) GENENTECH INC.
(GETH) GENENTECH INC.
(ery Match 100.0%; Score 713; DB 9; Length 713;
ADB21851 standard; CDNA, 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA.
US203082765-A1.
01-MAY-2003.
(GETH ) GENENTECH INC.
ery Match
st Local Similarity 100.0%; Score 713; DB 9; Length 713;
st Local Similarity 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                Length 713;
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(GETH ) GENENTECH INC.
ery Match 100.0%; Score 713; DB 9; Length 713;
ery Match 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082700-A1.
US2003082700-A1.
(GETH) GENENTECH INC.
ery Match
st. Local Similarity 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003054516-A1.
CO-MAR-2003.
(GETH ) GENENTECH INC.
ery Match
st Local Similarity 100.0%; Pred. No. 3.3e-155;
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Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003044806-A1.
06-MAR-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                   ADA87053 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 US2003082709-A1.
                                                                                                                                                                                                                              Score 713; DB 9; I
Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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820030924706-A1.
01-MAY-2003.
(GETH ) GENENTECH INC.
                                                                                                                                  Human PRO polynucleotide #237.
US2003068797-Al.
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01-MAY-2003.
(GETH ) GENENTECH INC.
100.0%;
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Best Local Similarity
RESULT 78
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Best Local Similarity
RESULT 76
                                                                                                         Best Local Similarity RESULT 72
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Best Local Similarity
RESULT 73
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Best Local Similarity
RESULT 75
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Best Local Similarity
RESULT 77
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RESULT 79
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                                                                                            Query Match
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Human cDNA encoding secreted/transmembrane polypeptide PRO1159 US2003040473-A1.
  Length 713;
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                                                                                                                                                                                      AUA88708 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003073213-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US20020097344-A1.
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US2003059782-A1.
27-MAR-2003.
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100.0%; Score 713; DB 9; 1
100.0%; Pred. No. 3.3e-155;
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Best Local Similarity 100.0%; Pred. No. 3.3e-155;
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Best Local Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
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US2003022239-A1.
                                                            CDNA; 713 BP.
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Length 713;
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(GETH) GENENTECH INC.
ery Match 100.0%; Score 713; DB 9; Length 713;
ery match 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US200308747-A1.
(GB-MAY-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                    Score 713; DB 9; Length 713;
Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA US2003082712-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082766-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082689-A1.
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Pred. No. 3.3e-155;
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100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Human PRO polynucleotide SEQ ID NO 473.
US2003077712-A1.
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01-MAY-2003.
(GETH ) GENENTECH INC.
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08-MAY-2003.
(GETH ) GENENTECH INC.
100.0%;
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01-MAY-2003.
(GETH ) GENENTECH INC.
""reft 100.0%; "
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US2003087352-Al.
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Human PRO polynucleotide #237.
US2003082698-A1.
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Human PRO polynucleotide #237.
US2003077711-A1.
24-APR-2003.
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(GETH ) GENENTECH INC.
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LAY MATCH

Best Local Similarity
RESULT 97
                                                                             GENENTECH INC
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Best Local Similarity
RESULT 92
ID ADA92450 standard; cf
DE Novel human secreted
PN US203082712-A1.
PD 01-MAY-2003.
PA (GETH ) GENENTECH INC
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Best Local Similarity
RESULT 99
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Best Local Similarity
RESULT 93
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Best Local Similarity
RESULT 94
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003082696-A1.
01-MAY-2003.
(CETH.) GENENTECH INC.
                                                                                                                        CDNA
                                                                                                                                                                                                                                                 ADB47221 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003082687-A1.
                                                                                                  AD539598 standard; cDNA; 713 BP.

Novel human secreted and transmembrane protein PRO1159
US203082764-A1.
01-MAY-2003.
(GEH) GENENTECH INC.
100.0%; Score 713; DB 10; Lengt
st Local Similarity 100.0%; Pred. No. 3.3e-155;
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(GETH) GENENTECH INC.

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24-APR-2003.
(GETH ) GENENTECH INC.
(ETY MATCh 100.0%; Score 713; DB 10;
ery Match 100.0%; Pred. No. 3.3e-155;
PD 01-MAY-2003.

PA (GETH ) GENENTECH INC.

Query Match

Best Local Similarity 100.0%; Pred. No. 3.3e-155;

RESULT 100
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(GETH ) GENENTECH INC.
(GETY Match 100.0%; Score 713; DB 10;
ery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Human PRO polymucleotide SEQ ID NO 473.
US2003077720-Al.
24-APR-2003.
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ID NO 473.
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Human PRO polynucleotide SEQ ID NO 473.
US2003077716-A1.
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ID NO 473.
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ID NO 473
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Human PRO polynucleotide #237.
US2003082697-A1.
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24-APR-2003.
(GETH) GENENTECH INC.
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Human PRO polynucleotide SEQ
US2003077717-A1.
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Human PRO polynucleotide SEQ
US2003077719-A1.
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Human PRO polynucleotide SEQ
US2003077718-A1.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 102
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Best Local Similarity
RESULT 104
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Best Local Similarity
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Ouery Match

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RESULT 119
ID ADCISSO47 standard; CDNA; 713 BP.
DB Novel human secreted and transmembrane protein CDNA Seq ID473
PN US2003087365-Al.
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Novel human secreted and transmembrane protein cDNA Seg ID473.
US2003087366-A1.
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Novel human secreted and transmembrane protein cDNA Seg ID473
US2003087364-A1.
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Novel human secreted and transmembrane protein cDNA Seq ID473
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Novel human secreted and transmembrane protein cDNA Seq ID473.
US2003087460-A1.
08-MAY-2003.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2002037367-A1.
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               ) 15-MAY-2003.

\( \text{GETH} \) \text{GENENTECH INC.} \)
\text{Query Match} \)
\text{Match} \)
\text{100.0$; Score 713; DB 10; \)
\text{Query.} \]
\text{100.0$; Pred. No. 3.3e-155; \)
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
(GETY MATCh 100.0%; Score 713; DB 10;
(ery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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08-May A.c.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 126
ID ADC59176 standard,
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ID ADC60592 standard;
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Best Local Similarity
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                                      Length 713;
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                                  Query Match

Best Local Similarity 100.0%; Pred. No. 3.3e-155;

RESULT 109

ID ADB46641 standard; CDNA; 713 BP.

DB Novel human secreted and transmembrane protein PRO1159 CDNA.

PN US2003082692-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003092107-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003092105-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003092106-A1.
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Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003049681-A1.
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(GETH ) GENENTECH INC.
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lery Match
100.0%; Score 713; DB 10;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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US2003064375-A1.
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Human PRO polynucleotide #118.
US2003045463-A1.
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15-MAY-2003.
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(GETH ) GENENTECH INC.
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RESULT 118
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Best Local Similarity
RESULT 111
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RESULT 112
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RESULT 113
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Best Local Similarity
RESULT 115
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Best Local Similarity
RESULT 116
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Best Local Similarity
RESULT 117
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Query Match

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ADD09575 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003194775-A1.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 138
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Best Local Similarity
RESULT 140
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RESULT 146
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Best Local Similarity
RESULT 139
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ID ADD06995 standard;
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                                                                     ADC58624 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein cDNA Seg ID473.
US2003087346-A1.
                                                                                                                     08-MAY-2003.
(GETH ) GENENTECH INC.
(ery Match initarity 100.0%; Score 713; DB 10; Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA, US2003082546-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003068623-A1.
10-APR-2003.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003092104-A1.
15-MAY-2003.
(GETH ) GENENTECH INC.
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003087348-A1.
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DE Human PRO polynucleotide #237.

PN US2003194773-A1.

PD 16-OCT-2003.

QUETY Match ) GENENTECH INC.

QUETY Match

Best Local Similarity 100.0%; Pred. No. 3.3e-155;

RESULT 136
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Pred. No. 3.3e-155;
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                      Score 713; DB 10;
Pred. No. 3.3e-155;
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ID ADC69709 standard; cDNA; 713 BP.

CDNA encoding human PRO polypeptide #237

PN US2003194770-A1.
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ID ADC82033 standard; CDNA; 713 BP.
DE Human PRO polynucleotide #118.
PN US2003083461-A1.
PD 01-MAY-2003.
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Query Match
Best Local Similarity 100.0%;
RESULT 128
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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...dry Match
Best Local Similarity RESULT 129
ID ADC14676 stand
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RESULT 130
ID ADD08208 stand
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Best Local Similarity

RESULT 131

ID ADD01298 stand*

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Best Local Similarity
RESULT 137
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Best Local Similarity
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(GETH ) GENENTECH INC.
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ery match 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003092103-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2001293300-A1.
ADD07675 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2002193299-A1.
(GETH ) GENENTECH INC.
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Novel human secreted and transmembrane protein PRO1159 cDNA 10/520030073090-A1.
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LETY MATCh 100.0%; Score 713; DB 10;
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PA (GETH ) GENENTECH INC.

QUERY MAtch 100.0%; Score 713; DB 10;

Query Match 100.0%; Pred. No. 3.3e-155;

RESULT 141
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155,
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Human PRO polynucleotide #237.
US2003194771-A1.
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08-MAY-2003.
(GETH ) GENENTECH INC.
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Length 713

CDNA

Length 713

Length 713;

RESULT 149

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Query Match 100.0%; Score 713; DB 10; Length 713; Best Local Similarity 100.0%; Pred. No. 3.3e-155; RESULT 160
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003087304-A1.
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                                                                                                                                                                                   Novel human secreted and transmembrane protein PRO1159 US2003203432-A1.
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(GETH ) GENENTECH INC.
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LETY MATCh 100.0%; Score 713; DB 10;
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(GETH ) GENENTECH INC,
lery Match 100.0%; Score 713; DB 10;
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(GETH ) GENENTECH INC.
sry Match
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(GETH ) GENENTECH INC.

Query Match

Best Local Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155,
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cDNA encoding human PRO polypeptide #237
US2003199056-A1.
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US2003199055-A1.
RESULT 156

ID ADD02108 standard; CDNA; 713 BP.
DE Human PRO polynucleotide #237.
PN US2003203430-A1.
PD 30-OCT-2003.
PA (GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADD92607 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003199030-A1.
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Human PRO polynucleotide #237.
US2003199057-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
Query Match 1
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003203438-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
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lery Match
100.0%; Pred. No. 3.3e-155;
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16-OCT-2003.
(GETH ) GENENTECH INC.
(ery Match 100.0%; Score 713; DB 10;
ery Match 100.0%; Pred. No. 3.3e-155;
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CDNA encoding human PRO polypeptide #237.
US2003194769-A1.
US2003194769-A1.
(GETH) GENENTECH INC.
ery Match
st Local Similarity 100.0%; Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
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16-OCT-2003.
(GETH ) GENENTECH INC.
100.0%; Score 713; DB 10;
Lery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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cDNA encoding human PRO polypeptide #237
US2003194792-A1.
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                                                                              Human PRO polynucleotide #118.
US2003059783-A1.
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Human PRO polymucleotide #118.
022003077594-A1.
24-APR-2003.
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30-OCT-2003.
(GETH ) GENENTECH INC.
100.0%;
                                                  Best Local Similarity RESULT 147
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Best Local Similarity
RESULT 155
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RESULT 148
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Best Local Similarity
RESULT 150
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Best Local Similarity
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Best Local Similarity
RESULT 152
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Best Local Similarity
RESULT 153
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Best Local Similarity
RESULT 154
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(GETH ) GENENTECH INC.

Query Match

Best Local Similarity 100.0%; Score 713; DB 10; Length 713;

RESULT 168

ID ADD92055 standard; cDNA; 713 BP.

DE Human PRO polynucleotide #237

PN US2001199053-A1.

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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003194791-A1.
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(GETH) GENENTECH INC.
100.0%; Score 713; DB 10;
107. Match 100.0%; Pred. No. 3.3e-155;
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                                                                     100.0%; Score 713; DB 10; 100.0%; Pred. No. 3.3e-155;
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cDNA encoding human PRO polypeptide #237.
US2003203428-A1.
                                                                                                                       ADE42106 standard; cDNA; 713 BP. Human PRO polymucleotide #237. US2003194772-A1. (GETH ) GENENTECH INC.
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Human PRO polynucleotide #237.
US2003194768-A1.
16-OCT-2003.
(GETH ) GENENTECH INC.
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RESULT 171
ID ADD80112 standar
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(GETH ) GENENTECH INC.
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                                                    (GETH ) GENENTECH INC
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Best Local Similarity
RESULT 167
ID ADE17923 standard; CDD
DE Human PRO polynucleot:
PN US2003199023-A1.
PD 23-OCT-2003.
PA (GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 169
ID ADE33518 standard; cD
DE Novel human secreted
PN US2003194767-A1.
PD 16-OCT-2003.
PA (GETH ) GENENTECH INC
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Best Local Similarity
RESULT 172
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Novel human secreted and transmembrane protein PRO1159 cDNA
US2003087305-A1.
08-MAY-2003.
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Novel human secreted and transmembrane protein PRO1159 US2003194766-A1.
USACCT-2003.
23-OCT-2003.
(GETH) GENENTECH INC.
(ery Match 100.0%; Score 713; DB 10; ery Match 100.0%; Pred. No. 3.3e-155;
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PA (GETH ) GENENTECH INC.
Query Match
Best Local Similarity 100.0%; Pred. No. 3.3e-155;
RESULT 181
                                                                                                                                                 Human PRO polynucleotide #237.
US2003199033-A1.
23-OCT-2003.
(GETH ) GENENTECH INC.
527 Score 713; DB 10; sty March
st Local Similarity 100.0%; Pred. No. 3.3e-155;
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Human PRO polynuclectide #237.
US200139902-A1.
23-OCT-2003.
(GETH ) GENENTECH INC.
CETY MAtch
St. Local Similarity 100.0%; Pred. No. 3.3e-155; St. Local Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155,
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cDNA encoding human PRO polypeptide #237
US2003203429-A1.
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cDNA encoding human PRO polypeptide #237.
US2003207418-A1.
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Human FRO polynucleotide #237.
US2003199028-A1.
23-OCT-2003.
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30-OCT-2003.
(GETH ) GENENTECH INC.
"a+ch 'la+ity 100.0%;
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                                                                                           Best Local Similarity
RESULT 175
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Best Local Similarity
RESULT 179
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RESULT 183
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2001049682-A1.
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Novel human secreted and transmembrane protein PRO1159 CDNA.
US2003201777-A1.
                                                                                                                                                                                                    ADH55955 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207379-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADI64174 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207385-A1.
06-NOV-2003.
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Novel human secreted and transmembrane protein PRO1159 CDNA.
US2003207386-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US200207388-A1.
                                                                            CDNA
                                        Novel human secreted and transmembrane protein PRO1159 052003201381-A1.
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(GETH ) GENENTECH INC.
Lery Match 113; DB 10;
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Pred. No. 3.3e-155;
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(GETH) GENENTECH INC.
ery Match 100.0%; Score 713; DB 10;
ery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
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ID AD13557 standard; cDNA; 713 BP.
DE Human PRO polynucleotide #118.
PN US2003050457-A1.
PD 13-MAR-2003.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 196
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Best Local Similarity
RESULT 202
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RESULT 197
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ID ADH82036 standard;
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(GETH) GENENTECH INC.
(ery Match 100.0%; Score 713; DB 10; Length 713;
(ery Match 100.0%; Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
ery Match imilarity 100.0%; Score 713; DB 10; Length 713;
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Novel human secreted and transmembrane protein PRO1159 cDNA US2003207355-A1.
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1D ADG23246 standard; cDNA; 713 BP.

DE Novel human secreted and transmembrane protein PRO1159 cDNA

US2003207384-A1.
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
ery Match
(ery Match 100.0%; Score 713; DB 10;
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16-OCT-2003.
(GETH ) GENENTECH INC.
(ETY Match 100.0%; Score 713; DB 10;
ETY Match 100.0%; Pred. No. 3.3e-155;
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PA (GETH ) GENENTECH INC.

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100.0%; Score 713; DB 10;

Best Local Similarity 100.0%; Pred. No. 3.3e-155;

RESULT 191
(GETH ) GENENTECH INC.

ry Match 100.0%; Score 713; DB 10;

+ Tocal Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Human PRO1159 nucleotide sequence SEQ ID NO:376.
US2001298148-Al.
26-DEC-2002.
(GETH ) GENENTECH INC.
100.0%; Score 713; DB 10;
St Local Similarity 100.0%; Pred. No. 3.3e-155
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                                                                          ADE40986 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003199031-A1.
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DE Human PRO polynucleotide #237.
PN US2003199034-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 192
ID AbG80111 standard;
                   Query Match
Best Local Similarity
RESULT 184
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RESULT 187
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Best Local Similarity
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RESULT 188
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AUNISSOO standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO11S9
10S2003083356-A1.
08-MAY-2003.
(GETH) GENENTECH INC.
ery Match 100.0%; Score 713; DB 11;
ery Match 100.0%; Pred. No. 3.3e-155;
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cDNA encoding human PRO polypeptide #237.
US2003092108-A1.
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US2003100087-A1.
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US2003203440-A1.
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Human PRO polynucleotide #237.
US2003092113-A1.
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Human PRO polynucleotide #237.
US2003211571-A1.
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08-WAY-2003.
(GETH ) GENENTECH INC.
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PA (GFTH ) GENENTECH INC.

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(GETH ) GENENTECH INC.
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JUEY MATCH
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Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003087353-A1.
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                                                                                                                                                                                                                                                       Human cDNA encoding secreted/transmembrane protein PRO1159.
US2003032023-A1.
                                                                                                                ABX80364 standard; DNA; 713 BP.
Novel human secreted or transmembrane protein PRO1124 DNA US2002132252-A1.
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Novel human secreted and transmembrane protein PRO1159
US2003087355-A1.
                                                                     100.0%; Score 713; DB 10; 100.0%; Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
Lery Match 100.0%; Score 713; DB 10;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
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    ABX77952 standard; cDNA; 713 B
Human PRO polynucleotide #118.
US2003027163-A1.
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13-FEB-2003.
(GETH ) GENENTECH INC.
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COETH ) GENENTECH II

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RESULT 207

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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 204

ID ACA69270 stand

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Best Local Similarity
RESULT 206
ID AEX90341 standard; CD
DE Human secreted/transm
PN US2002160384-A1.
PD 31-OCT-2002.
PA (GETH ) GENENTECH INC
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Best Local Similarity
RESULT 209
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Best Local Similarity
RESULT 205
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                                                                                    Local Similarity
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                                                      06-FEB-2003.
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ID ADN160
DE NCVE1
PN US2003
PD 08-MAY
PA (GETH
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PD 15-MAY-2003.
PA (GETH) GENENTECH INC.
Query Match 100.0%; Score 713; DB 12; Length 713; Best Local Similarity 100.0%; Pred. No. 3.3e-155; RESULT 217
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Length 713,
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                                                       ADN14948 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003087357-A1.
                                                                                                                                                                                         Novel human secreted and transmembrane protein PRO1159 (US2003092115-A1.
15-MAY-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
2ry Match 100.0%; Score 713; DB 12;
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PA (GETH ) GENENTECH INC.

Querry Match 100.0%; Score 713; DB 12;

Best Local Similarity 100.0%; Pred. No. 3.3e-155;

RESULT 219
Score 713; DB 11;
Pred. No. 3.3e-155,
                                                                                                                                                              Score 713; DB 11;
Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                            100.0%; Score 713; DB 12; 100.0%; Pred. No. 3.3e-155;
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US2003092110-A1.
15-MAY-2003.
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us-09-989-293a-376.rng.spdi

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Best Local Similarity 100.0%; Pred. No. 3.3e-155;
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Human PRO polynucleotide #237.
US2003207352-A1.
                                                                                                                                                                                                                                                                                                                          CDNA; 713 BP
                        RESULT 230
ID ADE93466 standard; CDNA; 713 B
DE Human PRO polynucleotide #237.
PN US2003199060-A1.
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                                                                                                23-OCT-2003.
(GETH ) GENENTECH INC.
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RESULT 235
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ID ADG20245 standard;
                                                                                                                                                 Local Similarity
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Best Local Similarity
RESULT 232
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                         ADE92362 standard;
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                              Length 713;
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(GETH ) GENENTECH INC.

2ry Match 100.0%; Score 713; DB 12;
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(GETH ) GENENTECH INC.
ery Match 100.0%; Score 713; DB 12;
ery Match 100.0%; Pred. No. 3.3e-155;
                                                                                                                         CST. CGT.H.) GENENTECH INC. (GET.H.) GENENTECH INC. 100.0%; Score 713; DB 12; ery Match 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                   ADE89336 standard; cDNA; 713 BP.

Human FRO polynucleotide #237.

U32003199062-A1.

U3-OCT-2003.

(GETH) GENEWIECH INC.

EY Match

St Local Similarity 100.0%; Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
(GETH ) GENENTECH INC.
ery Match
100.0%; Score 713; DB 12;
rimilarity 100.0%; Pred. No. 3.3e-155;
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ID ADB94804 standard; cDNA; 713 BP.
DB cDNA encoding human PRO polypeptide #237.
PP 23-OCT-2003.
PA (GETH ) GENENTECH INC.
                                                                            ADE24645 standard; cDNA; 713 BP.
cDNA encoding human PRO polypeptide #237
US2003092111-A1.
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                                                                                                                                                              Query Match
Best Local Similarity 100.0%; Pro
RESULT 222
ID ADD87470 standard; CDNA; 713 BP.
DE Human PRO polynucleotide #237.
PN US2003203439-A1.
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Human PRO polynucleotide #237.
US2003199054-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                            Query Match
Best Local Similarity
RESULT 221
                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 223
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Best Local Similarity
RESULT 224
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Best Local Similarity
RESULT 225
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Best Local Similarity
RESULT 228
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Best Local Similarity
RESULT 229
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100.0%; Score 713; DB 12; Length 713; 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003199058-A1.
                                                                                                                                                                                                                                                              Novel human secreted and transmembrane protein PRO1159 cDNA. US2003199051-A1.
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Novel human secreted and transmembrane protein PRO1159 US2003207360-A1.
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(GETH ) GENENTECH INC.
iry Match 100.0%; Score 713; DB 12;
rranal Similarity 100.0%; Pred. No. 3.3e-155;
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23-OCT-2003.
(GETH ) GENENTECH INC.
100.0%; Score 713; DB 12;
1ery Match 100.0%; Pred. No. 3.3e-155;
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Human PRO polynucleotide #237.
US200199663-Al.
23-OCT-2003.
(GETH ) GENENTECH INC.
ETY MAtch
100 0%; Score 713; DB 12;
ETY Match
100 0%; Pred. No. 3.3e-155;
Et Local Similarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
                                                             ADF35047 standard; cDNA; 713 BP.

OSDA encoding human PRO polypeptide #237.
23-0CT-2003.

(GERH) GENENTECH INC.

GERTH) GENENTECH INC.

INO.08; SCOTE 713; Struct Inc.
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025003207376-A1.
06-NOV-2003.
(GETH ) GENENTECH INC.
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Query Match
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                                                                                                                                                                                Novel human secreted and transmembrane protein PRO1159 cDNA US2003207426-A1.
                                                                                                                                                                                                                   PD 06-NOV-2003.
PA (GETH ) GENENTECH INC.
Querry Match 100.0%; Score 713; DB 12;
Best Local Similarity 100.0%; Pred. No. 3.3e-155;
RESULT 241
                                                                                                           Score 713; DB 12;
Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
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CDNA encoding human PRO polypeptide #237.
06-NOV-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADG19585 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237. US2003207425-A1.
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ID ADG03553 standard; CDNA; 713 BP. DE Human PRO Polynucleotide #237. PN 052003207351-A1. PD 06-NOV-2003. PA (GETH ) GENENTECH INC.
           Human PRO polynucleotide #237.
US2003207422-A1.
06-NOV-2003.
(GTH) GENEVIECH INC.
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                                                                                                                                                                                                                                                                                                             ADF98722 standard; cDNA; 713 BP.
Human PRO polynucleotide #237.
US2003208055-A1.
                                                                                                                                                                 ADG24368 standard; cDNA; 713 BP.
                                                                                                         100.0%;
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Best Local Similarity 100.0%;
RESULT 247
RESULT 239
ID ADF98151 standard; CDNA; 71
DE Human PRO polynucleotide #2
PN US2003207422-A1.
PD 06-NOV-2003.
PA (GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 246
ID ADG19585 standard; cDl
DE cDNA encoding human Pl
PUS2003207425-A1.
PD 06-NOV-2003.
PA (GETH ) GENEWTECH INC.
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Best Local Similarity
RESULT 244
                                                                                                                         Best Local Similarity RESULT 240
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ID ADG0531
DE Human P
PN US20032
PD 06-NOV-PA (GETH )
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Length 713;
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US2003207427-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADG23816 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US2002207389-A1.
ADGO8479 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207424-A1.
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Novel human secreted and transmembrane protein PRO1159
US2003207350-A1.
06-NOV-2003.
(GETH) GENENTECH INC.
100.0%; Score 713; DB 12; Lengt
Ext Local Similarity 100.0%; Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 US2003207356-A1.
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(GETH) GENENTECH INC.

Query Match

100.0%; Score 713; DB 12;

Query Match

100.0%; Pred. No. 3.3e-155;
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PA (GETH) GENENTECH INC.

Query Match

100 0%; Score 713; DB 12;

Best Local Similarity 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                             PD 06-NOV-2003.
PA (GETH ) GENENTECH INC.
Query Match
Best Local Similarity 100.0%; Pred. No. 3.3e-155;
RESULT 253
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(GENH ) GENENTECH INC.
Lery Match 100.0%; Score 713; DB 12;
Lery Match 100.0%; Pred. No. 3.3e-155;
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                                                                                   100.0%; Score 713; DB 12; 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
                                                                                                                                                    cDNA encoding human PRO polypeptide #237 US2003219885-A1.
                                                                                                                                                                                                                                                                     APP97047 standard; cDNA; 713 BP. Human PRO polynucleotide #237. US2003207371-A1.
                                                                                                                                      CDNA; 713 BP.
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Human PRO polynucleotide #237.
US2003207423-A1.
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06-NOV-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 257
ID ADG55150
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RESULT 256
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                                                                                                 Best Local Similarity
RESULT 249
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                                                                                                                                      ADG15649 standard;
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Length 713;

Length 713;

Query Match

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Length 713;

Length 713;

Length 713

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CDNA

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06-NOV-2003

Query Match

Query Match

06-NOV-2003

RESULT 264

Query Match

Best Loc RESULT 263

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AUG53694 standard; CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA.
US203207415-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207421-A1.
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ADHI2096 standard, CDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 CDNA.
US2003207419-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207416-A1.
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Human CDNA encoding secreted/transmembrane protein PRO1159.
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Novel human secreted and transmembrane protein PRO1159
US2003207414-A1.
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(GETH ) GENENTECH INC.
sry Match 100.0%; Score 713; DB 12;
oran eimilarity 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
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Pred. No. 3.3e-155
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155
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06-NOV-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
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     US2003207363-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA. US2003194778-A1.
                                                                                                                                                                                                                                                      ADG62118 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA.
US200207428-Al.
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Novel human secreted and transmembrane protein PRO1159 cDNA,
US2003207362-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207384-A1.
06-NOV-2003.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207368-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207365-A1.
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Novel human secreted and transmembrane protein PRO1159
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0GTH) GENENTECH INC.
100.0%; Score 713; DB 12;
ery Match
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(GETH) GENENTECH INC.

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100.0%; Score 713; DB 12;

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(GETH ) GENENTECH INC.
ery Match 100.0%; Score 713; DB 12;
ery Match 100.0%; Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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No. 3.3e-155;
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Pred. No. 3.3e-155;
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 260
                                                                                      Best Local Similarity
RESULT 258
ID ADG61014 standard;
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RESULT 262
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Best Local Similarity
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Query Match

RESULT 261

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Best Local Similarity 100.0%; Pred. No. 3.3e-155;
RESULT 285
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(GETH ) GENENTECH INC.
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Pred. No. 3.3e-155;
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Novel human secreted and transmembrane protein PRO1159 cDNA.
US2003207367-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                 ADG61566 standard; cDNA; 713 BP. Novel human secreted and transmembrane protein PRO1159 cDNA. US2003207429-A1.
                                                                       ADGS6454 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207366-A1.
                                                                                                                                                                        Novel human secreted and transmembrane protein PRO1159 cDNA, 052003207378-A1.
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Novel human secreted and transmembrane protein PR01159 cDNA
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Munan cDNA encoding secreted/transmembrane protein PRO1159.
US200324358-A1.
04-DEC-2003.
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Human cDNA encoding secreted/transmembrane protein PRO1159
US2003219856-A1.
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
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cDNA encoding human PRO polypeptide #237.
US2003207361-A1.
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Human PRO polynucleotide #237.
US2003022331-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity 1
RESULT 282
ID A0554798 standard; CDN
DE Novel human secreted a
PD 06-NOV-2003.
PD 06-NOV-2003.
PA (GETH ) GENENTECH INC.
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Best Local Similarity
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                                          Best_Local Similarity
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Best Local Similarity
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RESULT 279
ID ADG61r
DE Nove'
PN US'
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Length 713;
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Novel human secreted and transmembrane protein PRO1159 CDNA.
US2004009547-A1.
15-JAN-2004.
GETH ) GENENTECH INC.
100.0%; Score 713; DB 12; Length 713
st Local Similarity 100.0%; Pred. No. 3.3e-155;
ADG10005 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA US2004009548-A1.
                                                                                                                                                AD115476 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207382-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA US2003207349-A1.
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Novel human secreted and transmembrane protein PRO1159 cDNA.
1052004039164-A1.
26-FBB-2004.
(GETH ) GENENTECH INC.
100.0%; Score 713; DB 12; Length 713; Et Local Similarity 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                             ADI14808 standard; cDNA; 713 BP.
Novel human secreted and transmembrane protein PRO1159 cDNA
US2003207383-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 713; DB 12;
Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 713; DB 12;
Pred. No. 3.3e-155;
                                                                                            Score 713; DB 12;
Pred. No. 3.3e-155;
                                                                                                                                                                                                                                            Score 713; DB 12;
Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 713; DB 12;
Pred. No. 3.3e-155;
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Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   100.0%; Score 713; DB 12; 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADJ65701 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237. US2004038335-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             cDNA encoding human PRO polypeptide #237
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              CDNA; 713 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         CDNA; 713 BP.
                                US2004000...
15-JAN-2004.
(GETH ) GENENTECH INC.
100.0%; St
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human PRO polynucieotide #237. US2004038336-A1.
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06-NOV-2003.
(GETH ) GENENTECH INC.
Match 'Towity 100.0%;
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26-FEB-2004.
(GETH ) GENENTECH INC.
Watch '--ity 100.0%;
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06-NOV-2003.
(GETH ) GENENTECH INC.
100.0%;
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Query Match
PAGE TOCAL SIMILARITY 100.0%;
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity
RESULT 291
ID ADJ77579 standard; CI
DE Human PRO polynucleot
PN US2004038336-A1.
PD 26-FEB-2004.
PA (GETH ) GENENTECH INC
                                                                                          Query Match
Best Local Similarity
RESULT 286
                                                                                                                                                                                                                                                        Best Local Similarity
RESULT 287
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Best Local Similarity
RESULT 289
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADM27837 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              US200408333-A1.
11-MAR-2004.
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05-DEC-2002.
(ISIS-) ISIS INNOVATION LTD.
ery Match
10.9%; Score 77.6; DB 8; Length 2298;
ery Match 13.1%; Pred. No. 3.7e-08;
## WO200277216-A2.

10 03-OCT-2002.

A (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH.

PA (NOVS) NOVARTIS-ERFINDUNGEN VERW GES MBH.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human secreted protein 5' EST, SEQ ID NO: 28212.
EP1033401-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    #5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  11.6%; Score 83; DB 3; 97.6%; Pred. No. 1.1e-09;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity 100.0%; Pred. No. 1.2e-RESULT 305
ID ADC10755 standard; CDNA; 1281 BP.
DE Human CDNA from extracellular matrix gene 66
PN US2003059875-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABZ24041 standard; DNA; 1606 BP.
Human Dectin-1 polypeptide encoding DNA
WO200296945-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Mouse Dectin-1 polypeptide encoding DNA WO200296945-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABV73363 standard; cDNA; 138 BP.
LLR-J24-stalk peptide encoding cDNA.
WO200277216-A2.
                                                                                                                                                                                                                                                                                                                          Human polynucleotide SEQ ID NO 223. US2002042386-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAV42548 standard; cDNA; 2298 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ABZ24043 standard; DNA; 2298 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 10.9%;
73.1%;
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(ISIS-) ISIS INNOVATION LTD.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Mouse dectin-1 cDNA.
WO9828332-A2.
02-JUL-1998.
(TEXA ) UNIV TEXAS SYSTEM.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               .T 308
AAC24137 standard; cDNA;
                                                                                                                                                                                                                                                                                                                                                                                                                   (ROSE/) ROSEN C A.
(RUBE/) RUBEN S M.
(BARA/) BARASH S C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity RESULT 307
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Best Local Similarity
RESULT 310
                                                                                                                                                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                                                                                                                               RESULT 304
ID ABQ66733 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Local Similarity
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(GEST ) GENSET.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match
                                                                                                                Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AAK81282 standard; DNA; 5709 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36094.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAK81284 standard; DNA; 336 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:36096.
WO200157182-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADF60937 standard; cDNA; 1018 BP.
Human long Schering dendritic cell membrane protein 4 (SDCMP4) cDNA.
US2003162955-Al.
                                                                                                                                                    Score 713; DB 12; Length 713; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 5709
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 1018;
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                                                                                                                                                                                                                                                                                                                    Length 713;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Length 741;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 712;
                                                                                                                                                                                                                                                               USACCETO...
22-APR-2004.
(GETH) GENENTECH INC.
100.0%; Score 713; DB 12;
ery Match
-' "'milarity 100.0%; Pred. No. 3.3e-155;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               99.0%; Score 705.6; DB 4; 99.4%; Pred. No. 2.6e-153;
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06-JAN-1999.

(TAKED TAKEDA CHEM IND LTD.

(TAKE) TAKEDA CHEM IND LTD.

19.6%; Score 140; DB 2; I

ery Match 100.0%; Pred. No. 1.1e-22;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                98.3%; Score 701; DB 5; 99.9%; Pred. No. 1.9e-152;
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                                              ADM42561 standard; cDNA; 713 BP. cDNA encoding human PRO polypeptide #237. US2004058424-A1.
                                                                                                                                                                   Best Local Similarity 100.0%; Pred. No. RESULT 295
ID ADM28423 standard; cDNA; 713 BP.
DE CDNA encoding human PRO polypeptide #237
PN US2004077064-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAZ07533 standard; cDNA; 1018 BP.
Human SDCMP4 polypeptide encoding cDNA
W09947673-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABV73351 standard; cDNA; 1153 BP. LLR-J24-2 polypeptide encoding cDNA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAX01260 standard, cDNA, 741 BP.
Human DC3' protein coding sequence.
JP11001497-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CDNA; 712 BP.
                                                                                     US2004050.
25-MAR-2004.
(GETH ) GENENTECH INC.
100.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             (HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                      09-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (LEBE/) LEBECQUE S J E. (PHIL/) PHILLIPS J H.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  23-SEP-1999.
(SCHE ) SCHERING CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  (CHAL/) CHALUS L.
(QUAN/) QUAN A B.
(BATE/) BATES E B M.
(GCRM/) GORNAN D M.
(SAEL/) SAELAND S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        07-DEC-2000.
(GETH ) GENENTECH INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
RESULT 302
                                                                                                                                                                                                                                                                                                                  Ouery Match
Best Local Similarity
RESULT 296
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AAC91481 standard;
Human PRO1159 cDNA.
WO200073452-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 297
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AAS31409 standard; cDNA; 1281 BP.
Human cDNA encoding a novel extracellular matrix protein, Seq ID No 223.
WO200155368-A1.
Length 1153;
                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 1281;
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                                                                                                                                Length 1281;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match 19.6%; Score 140; DB 4; I ery Match 100.0%; Pred. No. 1.2e-22;
                                                                                                                                                                                                                                                                                               19.6%; Score 140; DB 6; 100.0%; Pred. No. 1.2e-22;
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PA (HUMA-) HUMAN GENOME SCI INC.
Query Match
Best Local Similarity 100.0%; Pred. No. 1.2e-22; RESULT 306
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  19.6%; Score 140; DB 8; 1 100.0%; Pred. No. 1.3e-22;
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Pred. No. 3.7e-08;
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Query Match
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Human juvenile hormone esterase binding protein homologue cDNA, SEQ:988.
WO200157188-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            6.5%; Score 46.4; DB 5; Length 335913; 45.4%; Pred. No. 1.7;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 5; Length 335913;
                                                                                                                                                                                                                                                                                                                    PD 03-JAN-2002.
PA (EPIG-) EPIGENOMICS AG.
Query Match 6.7%; Score 47.6; DB 6; Length 12590;
Best Local Similarity 45.6%; Pred. No. 0.46;
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match
forward for
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 44.4; DB 4; Length 5007; Pred. No. 2.1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               6.3%; Score 44.6; DB 4; Length 1204; 49.4%; Pred. No. 1.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 4; Length 1204;
                                                                                                                                                                                      Length 528;
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Pred. No. 0.96;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABN80067 standard, DNA, 7041 BP.
Human chemically modified disease associated gene SEQ ID
W0201200927-A2.
(BPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                               Human immune system associated gene SEQ ID NO: 1893.
WO200200928-A2.
RESULT 311

ID AAV42551 standard; CDNA; 528 BP.

DE Mouse dectin-1 extracellular domain cDNA.

PN WO982832-42.

PD 02-0UL-1998.

PA (TEXA) UNIV TEXAS SYSTEM.

Query Match

10.5%; Score 75.2; DB 2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             6.3%; Score 44.6; DE 49.4%; Pred. No. 1.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                6.5%; Score 46.4; DI 45.4%; Pred. No. 1.7;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Aai61371 standard; DNA; 335913 BP.
Soybean 240017 region G3, SEQ ID NO: 2.
W0200151627-A2.
19-JUL-2001.
(MONS ) MONSANTO CO.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Human ovarian cancer DNA marker #21276.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAI61372 standard; DNA; 335913 BP.
Soybean 240017 region G3, SEQ ID NO: 3.
WO200151627-A2.
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Human polynucleotide SEQ ID NO 2368.
WC200157190-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             AAK53107 standard; cDNA; 1204 BP.
Human polynucleotide SEQ ID NO 2636.
WO200157190-A2.
                                                                                                                                                                                                                                                                       ABL33920 standard; DNA; 12590 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADL63064 standard; DNA; 1675 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 6.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         6.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       19-JUL-2001.
(MONS ) MONSANTO CO.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 315
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Ouery Match
Best Local Similarity
RESULT 319
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 317
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Best_Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match
Best Local Similarity
RESULT 320
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 09-AUG-2001
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         09-AUG-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      09-AUG-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        RESULT 314
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          RESULT 316
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         RESULT 318
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PN
PD
PA
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Length 115863;
                                                                                                 Length 6497;
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03-JAN-2002.
(BPIG-) EPIGENOMICS AG.
5.9%; Score 42.4; DB 6; Length 5304;
                                                                                                                                                                                                                                                                                                                                                                                PD 24-OCT-2002.

PA (INCY-) INCYTE GENOMICS INC.

Query Match 6.2%; Score 44.2; DB 8; Length 3906;

Best Local Similarity 47.0%; Pred. No. 2.2;

RESULT 323
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       6.1%; Score 43.2; DB 6; Length 6509; 46.9%; Pred. No. 4.2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DB 11; Length 22693; 6.1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 8; Length 5216;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 110000;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length 319;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 319;
                                                                                                                                                                              Human soft tissue sarcoma-upregulated DNA - SEQ ID 2504.
MO2004048938-A2.
10-JUN - 2004.
(PROT -) PROTEIN DESIGN LABS INC.
6.2%; Score 44.4; DB 12; Length of Local Similarity 58.2%; Pred. No. 4.1;
ADN02625 standard; cDNA; 6497 BP.
Liver disease associated protein Obcll gene sequence.
NO2004029287-A2.
08-APR-2004.
(ORLD-) ORIDIS BIOMED FORSCHUNGS & ENTWICKLUNGS.
CORP. Match
St. Local Similarity 47.8%; Pred. No. 2.2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ABL34019 standard; DNA; 5304 BP.
Human immune system associated gene SEQ ID NO: 1992.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          200
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABZ36196 standard; cDNA; 5216 BP.
Human secretory polynucleotide SPTM SEQ ID NO 360
WO200283876-A2.
                                                                                                                                                                                                                                                                                                                                  Human secretory polynucleotide SPTM SEQ ID NO 141 W0200283876-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         D ADL38504 standard; DNA; 319 BP.

Human ovarian cancer DNA marker #12394.

N W0200170959-A2.

2.7-SEP-2001.

A (WILL-) MILENNIUM PREDICTIVE MEDICINE INC.

Query Match
Best Local Similarity 43.7%; Pred. No. 2.8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 AD 3373 standard; DNA; 319 BP.
Human ovarian cancer DNA marker #6115.
W0200170979-A2.
27-SEP-2001.
(MILL-) MILLERNIUM PREDICTIVE MEDICINE INC.
6.0%; Score 42.8; DB 5;
ery Match
st Local Similarity 43.7%; Pred. No. 2.8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           6.2%; Score 44; DB 6; 54.3%; Pred. No. 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 44.2; DE
Pred. No. 2.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADQ19685 standard; DNA; 115863 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human NRG2 gene fragment #1.
MO2003031594-A2.
                                                                                                                                                                                                                                                                                                                       ABZ35977 standard; cDNA; 3906 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     24-OCT-2002.
(INCY-) INCYTE GENOMICS INC.
lery Match 6.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DNA; 6509
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                  Best Local Similarity RESULT 321
                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 322
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Best Local Similarity
RESULT 325
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Best Local Similarity
RESULT 327
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 329
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 324
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL32227 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Query Match
                                                                                                     Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match
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                                                                                                                                                                                                                                                                 Query Match
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No.

55.4%; Pred.

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ABX48619 standard; cDNA; 397 BP.
Bovine EST associated with lactation/muscle/fat deposition #13784.
US2002137139-A1.
26-SEE-2002.
                                                                                                                                                                                                                                                                                              ABK28222 standard; DNA; 11394 BP.
DNA transcription associated complementary genomic DNA:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABL33054 standard; DNA; 6294 BP.

Human immune system associated gene SEQ ID NO: 1027,
202002001928-A2.
(FPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 5.8%; Score 41.2; DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DB 6;
                                                                                                                                                                                                                                                           DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human soft tissue sarcoma-upregulated DNA - SEO WO2004048938-A2.
                                                                                                                                                                                      Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                     ABQ99653 standard; DNA; 104644 BP.
Human MS4A7, MS4A5 and MS4A12 genomic sequence
WO200262946-A2.
                      Human prostate expression marker cDNA 56008.

MO200160860-A2.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.

ery Match

st Local Similarity 48.1%; Pred. No. 6.6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 41.2; DB Pred. No. 6.8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ΩB
                                                                                                                                                                                                             WOLDER
03-JAN-2002.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
5.8%; Score 41.4; D'
12; Atch
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       53.7%; Pred. No. 6.6;
                                                                                                                                                                                                                                                                                                                                                                                                     Score 41.4; | Pred. No. 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.8%; Score 41.4;
56.1%; Pred. No. 20;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | 10-JUN-2004.
| (PROT-) PROTEIN DESIGN LABS INC.
| ery Match | 5.8%; Score 41.2;
| cry Match | 5.8%; Pred. No. 9;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADL38009 standard, DNA, 348 BP.
Human ovarian cancer DNA marker #11899.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DNA; 8323 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ADQ25148 standard; DNA; 1533 BP
                                                                                                                                                                                                                                                                                                                                                    WOZULZZZ
06-DEC-2001.
(EPIG-) EPIGENOMICS AG.
5.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                              Best Local Similarity RESULT 340
                                                                                                                                                                     ABL32058 standard;
                                                                                                                                                                                                                                                           Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 342
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         15-AUG-2002.
(UYDU-) UNIV DUKE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 27-SEP-2001
                                                                                                                   Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAH19176 standard; cDNA; 1414 BP.
Human secreted protein-encoding gene 7 cDNA clone HPWAY46, SEQ ID NO:17.
WO200132910-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 1414;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 1414;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 1414;
                                                                                                                                                                                                                                  (HUMA-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match
5.9%; Score 41.8; DB 4; Length 1414;
                                                                                                                                                                                                                                                                                                                                                                                                                      DB 8; Length 1414;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 5265;
                                                                                                                                    Length 5858
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 7058
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     23-AUG-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 5.8%; Score 41.4; DB 5; Length 420;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human chemically pretreated gene sequence #67 strand WO200202806-A2.
                                                          Human immune system associated gene SEQ ID NO: 1285.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          HA-NOV-2002.
(HUMA-) HUMAN GENOME SCI INC.
6ry Match
5.9%; Score 41.8; DB 10;
5.9%; Pred. No. 6.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                OS-MAY-2003.
(HUMA-) HUMAN GENOME SCI INC.
6ry Match
5.9%; Score 41.8; DB 10;
5.9%; Pred. No. 6.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.9%; Score 41.8; DB 10; 49.3%; Pred. No. 6.4;
                                                                                    wcacc-

Wcaccol BPIGENOMICS AG.

(EPIG.) EPIGENOMICS AG.

6ry Match 5.9%; Score 42.4; DB 6;

cry Match 75.9%; Pred. No. 6.3;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            06-02-2001.
(EPIG-2001.
(EPIG-) EPIGENOMICS AG.
(ETY Match 5.8%; Score 41.6; DB 6;
(ETY Match 55.6%; Pred. No. 9.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 41.6; DB 6;
Pred. No. 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADC73667 standard; DNA; 1414 BP.
Human secreted protein-related DNA - SEQ ID 300
WO2003038063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADD37667 standard; cDNA; 1414 BP.
Human secreted protein encoding sequence #149.
WO200290526-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DNA transcription associated genomic DNA #147
WO200192555-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                  5.9%; Score 41.8; DI 49.3%; Pred. No. 6.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human prostate expression marker cDNA 8821.
W0200160860-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ADAS6266 standard, DNA; 1414 BP.
Gene encoding human secreted protein #445.
WO200210294-A2.
27-DEC-2002.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                         Human secreted protein encoding cDNA.
WO2002102993-A2.
27-DEC-2002.
                                                                                                                                                                                                                                                                                                                              ADA40093 standard; cDNA; 1414 BP
                                          ABL33312 standard; DNA; 5858 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABK40052 standard; DNA; 7058 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WOZUCZZZ
10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
*** MATCh '''*** 49.5%;
                                                                                                                                                                                                                                                                                                                                                                                                (HUMA-) HUMAN GENOME SCI INC.
Best Local Similarity RESULT 330
                                                                                                                                Query Match
Best Local Similarity
RESULT 331
                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 333
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Query Match
Best Local Similarity
RESULT 334
                                                                                                                                                                                                                                                                                     Best Local Similarity RESULT 332
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Best Local Similarity RESULT 338
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
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Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Local Similarity
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Best Local Si
RESULT 336
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                        Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
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Length 104644;

Length 348;

Length 348;

Length 1533

12;

Length 397;

.. 8

ID 7968

Length 11394;

9

DB

Length 8323;

Length 592;

5,

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ABX46069 standard; cDNA; 419 BP.
Bovine EST associated with lactation/muscle/fat deposition #11234.
US2002137139-A1.
                        ABMA1811 standard; CDNA; 337 BP.
Bovine EST associated with lactation/muscle/fat deposition #6976.
                                                                                                                                                                               ABN80169 standard; DNA; 6467 BP.

Human chemically modified disease associated gene SEQ ID NO 186.

W020020927-A2.

03-JAN-2002.

(RPIG-) EPIGENOMICS AG.
DB 6; Length 6294;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 8; Length 419;
                                                                                                                                                                                                                                                                                                                                                         Length 774;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               DB 6; Length 301;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 298;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.7%; Score 40.6; DB 6; Length 301;
48.5%; Pred. No. 8.8;
                                                                                                                                                                                                                                                                                                                                                                                                              71
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 298
                                                                                                                                                                                                                                                   Score 41; DB 6; Length 6467; Pred. No. 14;
                                                                                                                                             DB 8; Length 337; 7.3;
                                                                                                                                                                                                                                                                                                                                                                                        Colon tumour related determined cDNA sequence for contig
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          RESULT 353

ID ABL38485 standard; cDNA; 301 BP.

DE Human colon tumour antigen polynucleotide SEQ ID NO:2074.

PN W0201015388-2.

PD 20-DEC-2011.

PA (CORI.) CORIXA CORP.

5.7%; Score 40.6; DB 6; Length 3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ABL37552 standard; cDNA; 301 BP.
Human colon tumour antigen polynucleotide SEQ ID NO:1141
WO200196388-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 40.6; DB 4;
Pred. No. 8.8;
                                                                                                                                                                                                                                                                                                                                                            DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       DB 8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABZ33193 standard; cDNA; 298 BP.
Human colon tumour cDNA contig 71 SEQ ID NO:556
WO200283070-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Score 40.6; DE Pred. No. 9.4;
                                                                                                                                                                                                                                                                                                                                                          Score 40.8; DE Pred. No. 9.6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.7%; Score 40.6; DI
48.5%; Pred. No. 8.8;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             5.7%; Score 40.6; Di
48.5%; Pred. No. 8.8;
5.8%; Score 41.2; 1
52.3%; Pred. No. 12;
                                                                                                                                             Score 41;
Pred. No.
                                                                                                                                                                                                                                                                                     Frog embryonic gene sequence Q9925688. US2002081610-A1.
                                                                                                                                                                                                                                                   5.8%;
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                                                                                                                                             5.8%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.7%;
                                                                                                                                                                                                                                                                                                                               27-JUN-2002.
(UYRQ) UNIV ROCKEFELLER.
ery Match
5.7%;
st Local Similarity 38.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 26-SEP-2002.
BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                       (BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                                                                                                                                                                                                                                                                                                                                                                                       12-JUL-2001,
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              24-OCT-2002.
(CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                            Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match
Best Local Similarity
RESULT 356
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity RESULT 354
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (CORI-) CORIXA CORP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                              Query Match
Best Local Similarity
RESULT 352
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Best Local Similarity
                                                                                                                                             Query Match
Best Local Similarity
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Query Match
Best Local Similarity
RESULT 348
                                                                             26-SEP-2002
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                                                                                                                                                                       RESULT 349
                                                                                                                                                                                                                                                                             RESULT 350
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ID AB
DE BO
PN US
PD 26
PA (B
PA (M
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Human signal peptide containing protein HSPP-80 cDNA SEQ ID NO:214. WO200000610-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 6; Length 14429;
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                                                                                                                                                                                                                                                                                                                                                                                DB 4; Length 1765;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 6; Length 5228;
                                                                                                                                                                                                                                          DB 3; Length 1130;
                                                                                                     Length 588
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human anglogenesis associated polynucleotide SEQ ID NO 127 WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                            RESULT 359
ID ABL34242 standard, DNA, 14429 BP.
DE Human immune system associated gene SEQ ID NO: 2215.
PN W020020328-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human immune system associated gene SEQ ID NO: 1628 00200200208-A2. 003-JAM-2002. (EPIG-) EPIGENOMICS AG.
ABN62683 standard; cDNA; 588 BP.

Human cancer related polynucleotide SEQ ID NO 2650.

W0200214500-A2.

CHIR ) CHIRON CORP.

(HYSE-) HYSEQ INC.
                                                                                                     DB 6;
                                                                                                                                                                                                                                                                                                             552
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Human metastasis associated gene SEQ ID NO: 142.
WO200177376-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABS58028 standard; DNA; 5847 BP.
Human serum amyloid A4 (SAA4) gene segment #1
                                                                                                                                                                                                                                                                                             ; cDNA; 1765 BP. coding sequence SEQ ID NO:
                                                                                                                                                                                               WOZDUCCO

06-JZN-2000.

(INCY) INCYTE PHARM INC.

5.7%; Score 40.6; D

5.7%; Score 40.6; D
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03-JAN-2002.
03-JAN-2002.
08-JAN-2002.
S.78; Score 40.6; D
nery Match
5.78; Pred. No. 20;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Ouery Match
Best Local Similarity 45.5%; Pred. No. 26;
RESULT 362
ID ABL33655 standard; DNA; 5228 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 40.4; I
Pred. No. 19;
                                                                                                     5.7%; Score 40.6; I
48.5%; Pred. No. 10;
                                                                                                                                                                                                                                                                                                                                                                                5.7%; Score 40.6; 1
48.5%; Pred. No. 13;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 40.6; 1
Pred. No. 20;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  5.7%; Score 40.4; 153.9%; Pred. No. 18;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 40.4; Bred. No. 18;
                                                                                                                                                         AAZ98188 standard; cDNA; 1130 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABQ67097 standard; DNA; 14429 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DNA; 50000 BP
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PA (EPIG-) EPIGENOMICS AG.

Query Match 5.7%;

Best Local Similarity 46.2%;

RESULT 365
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (EPIG-) EPIGENOMICS AG.

12 Match

13 Local Similarity 47.6%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABLS6202 standard, DNA;
AMEPV genome fragment#4
WO200212526-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              14-FEB-2002.
(UYFL ) UNIV FLORIDA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Lery Match
Best Local Similarity
RESULT 364
ID ABL34589 "
                                                                                                                                                                                                                                        Query Match
Best Local Similarity
RESULT 358
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Best Local Similarity
RESULT 361
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                                                                                                                Best Local Similarity RESULT 357
                                                                                                                                                                                                                                                                                        AAH98698 standard;
Human EST-derived of
WO200154477-A2.
                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
                                                                                                                                                                                                                                                                                                                                              02-AUG-2001.
(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        13-JUN-2002
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
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                                                                                                         Query Match
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Query Match

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Length 1399;
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17;
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                                                                                                                       Length 1399;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADAS7756 standard; DNA; 1399 BP.
BAC fragment containing human secreted protein gene #455.
WO2002102994-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human chemically pretreated gene sequence #60 strand WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ADAS7757 standard; DNA; 1399 BP.
BAC fragment containing human secreted protein gene W02002102994-A2.
(HUMA-) HUMAN GENOME SCI INC.
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                                                                                                                                                                                           1326.
                                                                                                                                                                                                                                                                                                          ADC74694 standard; DNA; 1399 BP.
Human secreted protein-related DNA - SEQ ID 1327.
WO2003038063-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       5.6%; Score 40; DB 10;
52.4%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 10;
17;
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ry Match 5.6%; Score 40; DB 10;

t Local Similarity 52.4%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 10;
                                                                                                                                                                                                                                                              10;
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28;
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27;
                                                                                                                      5.6%; Score 40; DB 8; 52.4%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ä
                                                                                                                                                                 Human secreted protein-related DNA - SEQ ID WO2003038063-A2.
                                                                                                                                                                                                                                                            DB
17:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Human immune system associated gene SEQ ID WO200200928-A2.
     52.4%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 40;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                              08-MAY-2003.
(HUMA-) HUMAN GENOME SCI INC.
PTY MATCH 5.6%; Score 40;
St Local Similarity 52.4%; Pred. No.
                                                                                                                                                                                                                                                              40 Y
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.6%; Score 40; 52.4%; Pred. No.
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Pred. No.
                                                                                                                                                                                                                 MAX-2003.
(HUMA-) HUMAN GENOME SCI INC.
(ery Match
cery match
scimilarity 52.4%; Pred.
                                                   Human secreted protein related DNA, W0200210293-A2.
W7-DEC-2002.
(HUMA-) HUMAN GENOME SCI INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADD38147 standard; cDNA; 1399 BP.
cDNA clone in ATCC deposit #41.
WC200290526-A2.
14-NOV-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ADD38148 standard; cDNA; 1399 BP.
cDNA clone in ATCC deposit #42.
WO200290526-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABL32281 standard; DNA; 13511 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABK40038 standard; DNA; 16258 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  WOZDOZZE
10-JAN-2002.
(EPIG-) EPIGENOMICS AG.
-- Match ''reity 47.2%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (HUMA-) HUMAN GENOME SCI INC. ry Match 5.6%;
                                     ADA41620 standard; DNA; 1399
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  27-DEC-2002,
(HUMA-) HUMAN GENOME SCI INC.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Best Local Similarity RESULT 381
   Best Local Similarity
                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 376
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 380
                                                                                                                                        Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
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Best Local Similarity
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                     RESULT
                                                                                                                                                                                                                                                                                                                PERE
                                                                                                                                                                                                                                                                                          Plasmodium falciparum chromosome 2 related DNA sequence SEQ ID NO:303 WO200025728-A2.
ABL70396 standard; DNA; 7104 BP.
Chemically treated cell signalling DNA sequence complementary to#143.
WO200202807-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAL03269 standard; DNA; 1399 BP.
Human reproductive system related antigen DNA SEQ ID NO: 5957.
WO200155320-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 6062;
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                                                                                     Length 7104;
                                                                                                                                                                                                                                                                                                                                                                                                                  DB 3; Length 4041;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 6561;
                                                                                                                                                                                                                          Length 3197
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABK28222 standard; DNA; 11394 BP.
DNA transcription associated complementary genomic DNA #48.
WO200192565-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Length 1399;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.6%; Score 40; DB 4; Length 1399; 52.4%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                               AAS61094 standard; DNA; 6062 BP.
Human gene regulation-associated gene oligonucleotide #49
WO200177375-A2.
                                                                                                                                  ACF05254 standard; cDNA; 3197 BP.
Plasmoddium falciparum cGMP dependent protein kinase cDNA, W02003054157-A2.
03-JUL-2003.
(MRRI) MERCK & CO INC.
15.6%; Score 40.2; DB 9; Length 3 t Local Similarity 44.5%; Pred. No. 18;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         gene SEQ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MCZCT-2001.
(EPGCT-2001.
(EPGG-) EPIGENOMICS AG.
(EPTG MATCH 5.6%; Score 40.2; DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Www.vor...
(BD13-JAN-2002.
(BP16-) EPIGENOMICS AG.
Ery Match
Ery Match
5.6%; Score 40.2; DB 6;
                                                                                  5.7%; Score 40.4; DB 6; 46.2%; Pred. No. 19;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  RESULT 369
ID ABN80020 standard; DNA; 6561 BP.
DE Human chemically modified disease associated
PN W020020027-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                 Score 40.2; Pred. No. 19;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 40.2; |
Pred. No. 23;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             02-AUG-2001.
(HUMA-) HUMAN GENOME SCI INC.
ery Match 5.6%; Score 40;
ery match 5.2.4%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADA41621 standard, DNA; 1399 BP.
Human secreted protein related DNA
WO2002102993-A2.
27-DEC-2002.
                                                                                                                                                                                                                                                                          AAA70170 standard; DNA; 4041 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAL03268 standard; DNA; 1399 BP
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06-DEC-2001.
(EPIG-) EPIGENOMICS AG.
5.6%;
                                                                                                                                                                                                                                                                                                                                                                                                               5.6%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 (HUMA-) HUMAN GENOME SCI INC
                                                                    EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                           11-MAY-2000.
(HOFF/) HOFFMAN S.
(CARUC) CARUCCI D.
(GARD/) GARDNER M.
(VENT/) VENTER J C.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity RESULT 373
                                                                                                                                                                                                                        Query Match
Best Local Similarity
RESULT 367
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 372
ID AAL03269 standard;
                                                                                                    Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
                                                  10-JAN-2002
(EPIG-) EPI
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Query Match

RESULT 368

Query Match

Query Match

Ouery Match

Query Match

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Query Match
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         ABL/10376 standard; DNA; 16258 BP.
Chemically treated cell signalling DNA sequence complementary to#133.
WOZO0202807-A2.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 39.8; DB 10; Length 16579; Pred. No. 31;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 39.8; DB 10; Length 16579;
Pred. No. 31;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 16579;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human chemically treated TPBF nucleotide sequence SEQ ID NO:18 WO2003072820-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Drosophila melanogaster genomic polynucleotide SEQ ID NO 34111 W0200111042-A2. 27-SEP-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 16579
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ADE37762 standard; DNA; 16579 BP.

Human chemically treated TPEF nucleotide sequence SEQ ID NO:8.
WO203072820-A2.
04-SEP-2003.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                DB 10; Length 1940;
                                                                                                                                                                                                                                                                                                              Length 1131;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 4; Length 2669;
                                                                                                                                                                                                   Length 874;
                                                                                 Length 16258
                                                                                                                                     Human breast cancer expressed polynucleotide 16595.
W0200151628-A2.
19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Ery Match
St. (Score 39.8) DB 4; Lest Local Similarity 46.0%; Pred. No. 17;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 10;
                                                                                                                                                                                                                                                                                                              DB 3;
                                                                              Query Match 5.6%; Score 40; DB 6;
Best Local Similarity 47.2%; Pred. No. 28;
                                                                                                                                                                                                                                                                           WACKULLER-2000.
(HUMA-) HUMAN GENOME SCI INC.
(HUMA-) HUMAN GENOME SCI INC.
16ry Match
1 "Amilarity 52.8%; Fred. No. 18;
                                                                                                                                                                                                                                                                                                                                                                                            5.6%; Score 39.8; I 49.8%; Pred. No. 21;
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Pred. No. 31;
                                                                                                                                                                                                                                         AACS9161 standard; cDNA; 1131 BP.
Human secreted protein cDNA sequence #5.
WO200055201-A1.
                                                                                                                                                                                                                                                                                                                                                       ABX77226 standard; DNA; 1940 BP.
DNA sequence of BAC clone /ctb-137N13.
WO200283897-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADB54245 standard; DNA; 16579 BP.
Pretreated genomic DNA region 169,
0002003072821-A2.
04-SEP-2003.
(BPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADB54117 standard; DNA; 16579 BP.
Pretreated genomic DNA region 41.
WO2003072821-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ADE37772 standard; DNA; 16579 BP.
                                                                                                                           AAL24138 standard; cDNA; 874 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ABL27546 standard; DNA; 2669 BP.
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04-SEP-2003.
(EPIG-) EPIGENOMICS AG.
5.6%;
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04-SEP-2003.
(EPIG-) EPIGENOMICS AG.
TW MAtch -- 1 arity 52.8%;
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Best Local Similarity
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RESULT 385
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RESULT 390
RESULT 383
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Human multidrug resistance associated protein gene associated sequence. WO200257410-A2.
25-JUL-2002.
(DNAS.) DNA SCI LAB INC.
5.6%; Score 39.8; DB 6; Length 98472; st Local Similarity 47.7%; Pred. No. 45;
                                                                                                                                                                                                                                                                                                                                                      AAK71165 standard; DNA; 20420 BP.
Human immune/haematopoietic antigen genomic sequence SEQ ID NO:27977.
WO200157182-A2.
AAS46313 standard; DNA; 16750 BP.
Tumour suppressor gene derived chemically modified sequence #35.
WO200168912-A2.
20-SEP-2001.
(EPIG-) EPIGENOMICS AG.
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, (EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
Query Match 5.6%; Score 39.8; DB 10; Length 98472;
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                                                                                                            5.6%; Score 39.8; DB 4; Length 16750; 52.8%; Pred. No. 31;
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Cancer based on CYP3A5 related polynucleotide SEQ ID NO:673
WO2003013534-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ADB20860 standard; DNA; 98472 BP.
MRP1 based cancer related nucleic acid SEQ ID NO:673.
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Human secreted protein gene 68 genomic DNA fragment WO200226931-A2.
                                                                                                                                                                                                   493
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MSI-H carcinoma genomic DNA sequence SEQ ID NO:40.
RX2004008012-A.
RX2004008012-A.
(KIMH/) KIM H G.
(KIMH/) KIM N G.
(LEEJ/) LEE J S.
(RHEE/) RHEE H S.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 6;
                                                                                                                                                                                                   Human immune system associated gene SEQ ID NO: WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                           WAGGE 2001.
(HUMA-) HUMAN GENOME SCI INC.
S.6%; Score 39.8; D
HERY MATCH 46.5%; Pred. No. 33;
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(HUMA-) HUMAN GENOME SCI INC.

5.6%; Score 39.8; D

ery Match

ciry Match

46.5%; Pred. No. 33;
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67.5%; Pred. No. 43;
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                                                                                                                                                                                                                                                                                      5.6%; Score 39.8; 1
52.8%; Pred. No. 31;
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Human UGT1A1 gene sequence SEQ ID NO:673.
WO2003013536-A2.
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20-FEB-2003.
(EPID-) BPIDAUROS BIOTECHNOLOGIE AG.
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(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
                                                                                                                                                                              DNA; 16750 BP
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(EPIG-) EPIGENOMICS AG.
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Best Local Similarity
RESULT 398
                                                                                                                Query Match
Best Local Similarity
                                                                                                                                                       RESULT 393
ID ABL32520 standard;
                                                                                                                                                                                                                                                                                                        Best Local Similarity RESULT 394
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US5116965-A
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                                                                                                      Ouery Match
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AACS8017 standard; DNA; 20674 BP.
Arachidonic acid metabolism related genomic biallelic marker #651
WO200047771-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5.6%; Score 39.8; DB 10; Length 243072; 52.8%; Pred. No. 55;
                                                                                                                                                                                                                                                                                                                                                            DB 10; Length 110000;
                                                                                                                      DB 10; Length 98472;
                                                                                                                                                                                                                                                         Length 98472;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABL33702 standard; DNA; 18624 BP.
Human immune system associated gene SEQ ID NO: 1675.
WO200200928-A2.
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(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.

5.6%; Score 39.8; DB 10;

ery Match

7.7%; Pred. No. 45;
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Photorhabdus luminescens nucleotide sequence #35
WO200294867-A2.
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56.0%; Pred. No. 36;
                                                                                                                                                                                                                                                                                                            DB 3;
                               ADB96932 standard; DNA; 98472 BP.
Human MDR1 related DNA sequence SEQ ID NO:673.
WO2003013537-A2.
                                                                                                                                                                      DNA; 98472 BP.
DNA sequence SEQ ID NO:673.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABA93723 standard; cDNA; 1464 BP.
Human amygdala derived cDNA clone amy2_24k15.
W0200198454-A2.
(GEHU-) GERMAN HUMAN GENOME PROJECT.
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28-SEP-2000.
(GEMY) GENETICS INST INC.
5.6%; Score 39.6; D
ery Match 56.0%; Pred. No. 44;
                                                                                                                    5.6%; Score 39.8; I
47.7%; Pred. No. 45;
                                                                                                                                                                                                                                                                                                          Score 39.8; 1
Pred. No. 46;
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Pred. No. 21;
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Pred. No. 36;
                                                                                                                                                                                                                                                                                                                                                          Score 39.8; Pred. No. 46;
47.7%; Pred. No. 45;
                                                                                 20-FEB-2003.
(EPID-) EPIDAUROS BIOTECHNOLOGIE AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               AA189341 standard; cDNA; 288 BP.
Human polynucleotide SEQ ID NO 9401.
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(CNRS ) CNRS CENT NAT RECH SCI.
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(EPIG-) EPIGENOMICS AG.
IY MATCh 5.6%;
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                                                                                                                                                                                                                                                                                                                                                          52.8%;
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WO200164835-A2.
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RESULT 408
                                                                                                                                                                    ADB92123 standard;
Human MDR1 related
WO2003013535-A2.
Best Local Similarity
RESULT 401
                                                                                                                    Query Match
Best Local Similarity
RESULT 402
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Best Local Similarity
RESULT 411
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Best Local Similarity
RESULT 404
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Best Local Similarity
RESULT 403
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Best Local Similarity
RESULT 405
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(HYSE-) HYSEQ INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         17-AUG-2000.
(GEST ) GENSET.
                                                                                                                                                                                                                                                                                                                                                                                                                                                          28-NOV-2002
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AAQ25532 standard; DNA; 1648 BP. Sequence of genomic clone contg. the entire Histidine-rich protein (HisRP) gene.
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                                                                                                                                                                                                  Length 1648
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  MBN80238 standard; DNA; 15121 BP.
Human chemically modified disease associated gene SEQ ID NO
WO200200927-A2.
                                                                                                                                                                                                                                                                                                                                  Human soft tissue sarcoma-upregulated DNA - SEQ ID 5934 WO200048938-A2.
10-UIN-2004.
(PROT-) PROTEIN DESIGN LABS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Signal transduction associated gene modified DNA #192.
WO20020020926-A2.
03-JAN-2002.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human chemically pretreated gene sequence #23 strand WO200202806-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human immune system associated gene SEQ ID NO: 1290 WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Human immune system associated gene SEQ ID NO: 1971
02020200928-A2.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 12;
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                                                                                                                                                                                                     DB 2;
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C. albicans BAX-associated cDNA fragment SEQ ID
WO200264766-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              В
                                                                                                                      SG-MAY-1992.
(SLOK ) SLOAN KETTERING INST CANCER.
SLOAN SLOAN KETTERING INST CANCER.
S-5*; Score 39.4; D
S-7-10 Similarity 52.1*; Pred. No. 24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Mozecz---
(33-JAN-2002.
(33-JAN-2002.
(BDIG-) BPIGENOMICS AG.
EPY Match
Fry 
                                                                                                                                                                                                                                                                                                                                                                                                                                                                             5.5%; Score 39.4; I
46.8%; Pred. No. 31;
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49.3%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 39.4;
Pred. No. 37;
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Pred. No. 38;
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Pred. No. 54;
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Pred. No. 57;
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Pred. No. 57;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Score 39.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ABL33317 standard; DNA; 13574 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ABL33998 standard; DNA; 21537 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ABK39964 standard; DNA; 15479 BP
                                                                                                                                                                                                                                                                                                   ADQ23114 standard; DNA; 5798 BP.
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03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
6-rv Match 5.5%;
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(EPIG-) EPIGENOMICS AG.
1ry Match 5.5%;
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17 Match 5.5%;

It Local Similarity 48.4%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Local Similarity
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RESULT 419
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Length 467;

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CDNA; 672 BP
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03-JDN-2002.
(EPIG-) EPIGENOMICS AG.
5.5%;
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  5.5%;
                                                                                                                                                                                                                                                                              5.5%;
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(EPIG-) EPIGENOMICS AG.
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                                                                                                                                                                                                                                                                                                                                                                          (CHIR ) CHIRON CORP. (HYSE-) HYSEQ INC.
Query Match
Best Local Similarity
RESULT 437
                                                                                                                                      Query Match
Best Local Similarity
RESULT 438
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Best Local Similarity
RESULT 439
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Best Local Similarity
RESULT 444
                                                                                                                                                                                                                                                                                                                      AAZ17651 standard;
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                                                                                                                             DB 6; Length 6045;
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                 Length 6045;
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
5.5%; Score 39; DB 5; Length 467;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 309;
                                                      Chemically treated cell signalling DNA sequence#257. Chemically treated cell signalling DNA sequence#257. W0200202807-A2.

10-JAN-2002.

SEGON BERGENOMICS AG.

5.5%; Score 39.2; DB 6; Lerent Local Similarity 52.4%; Pred. No. 35;
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                                                                                                                                                                                                                                                                                                                           DB 6;
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                 DB 6;
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ery March

7.5%; Score 39; DB 5;

rocal Similarity 48.1%; Pred. No. 21;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  nov44994 standard; cDNA; 309 BP.
Human prostate expression marker cDNA 44985.
WO20160860-A2.
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                                                                                                                                                                                                                                                                                 39.2; D
No. 64;
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No. 64;
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Pred. No. 38;
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Pred. No. 64;
                 Score 39.2;
Pred. No. 35;
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Pred. No. 64;
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Pred. No. 64;
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Pred. No. 73;
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                                                                                                                                                                                                                                                                                                                         Score 39.2;
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Human ovarian cancer DNA marker #11145.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ADI72106 standard; DNA; 467 BP.
Human ovarian cancer DNA marker #4848.
WC200170979-A2.
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Pred.
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                                                                                                                                                                                Carbamoyl-phosphate-synthetase II WO9412643-A1.
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                                                                                                                                                                      AAQ62924 standard; cDNA; 8920 BP.
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11-APR-2GO2.
(GENE-) GENE LOGIC INC.
5.5%;
                 Query Match 5.5%;
Best Local Similarity 52.4%;
RESULT 423
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                                                                                                                                                                                                                                                                                                                                      64.1%;
     (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                             (UNIX ) UNISEARCH LTD.
                                                                                                                                      Best Local Similarity RESULT 424
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Best Local Similarity
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Query Match
Best Local Similarity
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Best Local Similarity
RESULT 431
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Best Local Similarity
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Best Local Similarity
RESULT 426
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Best Local Similarity
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                                                                                                                             Query Match
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RESULT 425
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ADM91120 standard; DNA; 615 BP.
Human DNA encoding a pharmaceutically useful protein SeqID 513.
W02004020595-A2.
11-MAR-2004.
11-MAR-2004.
(RIVE-) FIVE RIME THERAPEUTICS INC.
(RIKE-) RIKEN INST PHYSICAL & CHEM RES.
(DNAF-) DNAFORM KK.
                                                ADM90702 standard; DNA; 615 BP.
Human DNA encoding a pharmaceutically useful protein SeqID 95.
WO2004020595-A2.
11-MAR-2004.
(FIVE-) FIVE PRIME THERAPEUTICS INC.
(RIKE-) RIKEN INST PHYSICAL & CHEM RES.
(NAR-) DNAFORM KK.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length 3629
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Length 7450;
                                                                                                                                                                     Length 615;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 1182;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 5195;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 6059;
                                                                                                                                                                                                                                                                                                                                             Length 615
                                                                                                                                                                                                                                                                                                                                                                                                                Human gene expression product cDNA sequence SEQ ID NO:5124 WO9938972-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Length 672;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      894
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ADI22698 standard, DNA, 3629 BP.
Human liver differentially expressed cDNA seq id
US2003165854-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DB 10;
35;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     DB 6;
40;
                                                                                                                                                                                                                                                                                                                                               12;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             4;
                                                                                                                                                                       5.5%; Score 39; DB 12; 59.5%; Pred. No. 24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    5.5%; Score 39; DB 2; 50.6%; Pred. No. 24;
Score 39; DB 5;
Pred. No. 23;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Human immune system associated gene SEQ ID NO: WO200200928-A2.
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genomic DNA #3.
                                                                                                                                                                                                                                                                                                                                               DB
24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB
27;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB
37;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Score 39; DB
Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Score 39;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 39;
Pred. No.
                                                                                                                                                                                                                                                                                                                                               Score 39;
Pred. No.
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Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             5.5%; Score 39; 50.6%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAKS2123 standard; cDNA; 1182 BP.
Human polynucleotide SEQ ID NO 668.
WQ200157190-A2.
09-AUG-2001.
(HYSE-) HYSEQ INC.
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DB 6; Length 5641;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Human immune system associated gene SEO ID NO: W0200200928-A2. W0200720028-A2. (EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ABX28257 standard; DNA; 8170 BP.
DNA transcription associated genomic DNA #66.
WO200192565-A2.
                                                                                                                                                                                           Query Match 5.4%; Score 38.8; I Best Local Similarity 58.8%; Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          OS-LOCATOR SAG.
Sry Match
St. Match

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     5.4%; Score 38.8; I
46.9%; Pred. No. 42;
                                                                                                                                                                                                                                                                                                                                                                                                                                             Score 38.8; I
Pred. No. 43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
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43.6%; Pred. No. 29;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.4%; Score 38.8; 56.2%; Pred. No. 48;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.4%; Score 38.6; 53.8%; Pred. No. 25;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         5.4%; Score 38.6; I
44.4%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Cotton expressed sequence tag, EST, #470.
US2004123338-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ADL37605 standard; DNA; 576 BP.
Human ovarian cancer DNA marker #11495.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        ADI72466 standard; DNA; 576 BP.
Human ovarian cancer DNA marker #5208.
27-SBP-2001.
                                                                                     DNA; 5893 BP.
                                                                                                                                                                                                                                                                                                 ADJ76057 standard; DNA; 5893 BP.
Marker gene SEQ ID NO:1309.
EP1394274-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ABL32062 standard; DNA; 9905 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   DNA; 2279 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                    Best Local Similarity 58.8%;
RESULT 456
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     AAQ49780 standard; DNA; 4526
                                                                                                         Marker gene SEQ ID NO:1207.
EP1394274-A2.
                                                                                                                                                                 03-MAR-2004.
(GENO-) GENOX RES INC.
                                                                                                                                                                                                                                                                                                                                                                                    03-MAR-2004.
(GENO-) GENOX RES INC.
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     Query Match
Best Local Similarity
RESULT 454
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Best Local Similarity RESULT 457
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Best Local Similarity
RESULT 462
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                                                                               ADJ75955 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ABL18982 standard;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            (PEKE ) PE CORP NY.
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WO9319079-A1.
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RESULT 458
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                                                                                                                                                                                ABL/0480 standard; DNA; 9742 BP.
Chemically treated cell signalling DNA sequence complementary to#185.
W0200202807-A2.
10-JAN-2002.
(RPIG-) EPIGENOMICS AG.
iv Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABX39417 standard; cDNA; 393 BP.
Bovine EST associated with lactation/muscle/fat deposition #4582.
US2002137139-A1.
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(FIVE-) FIVE PRIME THERAPEUTICS INC.

(FIVE-) FIVE PRIME THERAPEUTICS INC.

5.4%; Score 38.8; DB 12; Length 3470;

cry Match

58.8%; Pred. No. 38;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             5.4%; Score 38.8; DB 12; Length 5316; 42.6%; Pred. No. 42;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human presynaptic cytomatrix protein, PCLO, genomic sequence WO2004044164-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 89500
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           5.4%; Score 38.8; DB 8; Length 199; 54.1%; Pred. No. 21;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 8; Length 393;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Length 263
                                                                                                                                     Length 9180;
                                                                                                                                                                                                                                                                                                                                                             Length 9742
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Human immune system associated gene SEQ ID NO: 1369.
03-JAN-2002.
ABL33964 standard, DNA, 9180 BP.
Human immune system associated gene SEQ ID NO: 1937.
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Murine gene trapped sequence (GTS) SEQ ID No 981.
US2002161207-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DB 5;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         12;
                                                                                                                                     DB 6;
42;
                                                                                                                                                                                                                                                                                                                                                             DB 6;
43;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 7
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     5.4%; Score 38.8; I
46.1%; Pred. No. 24;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   5.4%; Score 38.8; 154.1%; Pred. No. 22;
                                                                                                                                  5.5%; Score 39; 56.7%; Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Score 39;
Pred. No.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ADL44393 standard; DNA; 263 BP.
Human ovarian cancer DNA marker #18283.
                                                                                                                                                                                                                                                                                                                                                             Score 39;
Pred. No.
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Human PRO polynucleotide #438.
WO2004043361-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                          ADO56275 standard; DNA; 89500 BP.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ADO36005 standard; DNA; 3470 BP.
Novel mouse gene sequence #678.
WO2004046310-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABX91637 standard; cDNA; 199 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      5.5%;
                                                                                                                                                                                                                                                                                                                                                          5.5%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             26-SEP-2002.
(BYAT/) BYATT J C.
(MATH/) MATHIALAGAN N.
(TAON/) TAO N.
(WARR/) WARREN W C.
                                                                            03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            31-OCT-2002.
(FRIE/) FRIEDRICH G.
(ZAMB/) ZAMBROWICZ B.
(SAND/) SANDS A.T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       27-MAY-2004.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (EPIG-) EPIGENOMICS AG
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RESULT 453
                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity RESULT 447
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 449
                                                                                                                                                            Local Similarity
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                                                                                                                                     Query Match
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DB 12; Length 5893;
                                                                                                                                                          DB 12; Length 5893;
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                                                                                                                                                                                                                                                                                                                       DB 6; Length 8170;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    DB 6; Length 9905;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 12; Length 256;
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(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC. 5 4*: Score 38.6; DB 5; Length 576;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      DB 4; Length 2279;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 576;
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Query Match
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                                                                                                                                                                                                                                                                                                     5.4%; Score 38.6; DB 6; Length 5376; 47.7%; Pred. No. 47;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Length 5962;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DB 6; Length 6591;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length 6015;
                                                                                            DB 2; Length 4526;
                                                                                                                                                                                                DB 2; Length 4526;
                                                                                                                                                                                                                                                                                                                                                                                                             Length 5416;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 6486;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Human angiogenesis associated polynucleotide SEQ ID NO WO200246454-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ABL33250 standard; DNA; 6591 BP.
Human immune system associated gene SEQ ID NO: 1223.
WO20020928-A2.
(BPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                           ABL33796 standard; DNA; 5416 BP.
Human immune system associated gene SEQ ID NO: 1769.
WO200200928-A2.
03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                                                                                                                                                                                                ABL33287 standard; DNA; 5962 BP.
Human immune system associated gene SEQ ID NO: 1260.
WO200200928-A2.
                                                                                                                                                                                                                                    ABL34150 standard; DNA; 5376 BP.
Human immune system associated gene SEQ ID NO: 2123
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     650
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(BPIG-) EPIGENOMICS AG.
Serv Match 5-4%; Score 38.6; DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                             DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Human immune system associated gene SEQ ID NO: W0200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ABL32583 standard; DNA; 13376 BP.
Human immune system associated gene SEQ ID NO:
WO200200928-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Score 38.6; I
Pred. No. 57;
                                                                                           5.4%; Score 38.6; 1
48.4%; Pred. No. 45;
                                                                                                                                                                                                                                                                                                                                                                                                            Score 38.6; 1
Pred. No. 47;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Score 38.6; Pred. No. 49;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 5.4%; Score 38.6;
44.1%; Pred. No. 49;
                                                                                                                                                                                                   4%; Score 38.6;
4%; Pred. No. 45;
    48.4%; Pred. No. 45;
                                                                (UNIW ) UNIV WASHINGTON STATE RES FOUND.
                                                                                                                                                                       16-MAR-1999.
(UNIW ) UNIV WASHINGTON STATE RES FOUND.
                            AAX06993 standard; cDNA; 4526 BP.
                                                                                                                                AAX19122 standard; DNA; 4526 BP. Tomato prosystemin genomic DNA. US5883076-A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ABQ67049 standard; DNA; 6486 BP.
                                        Tomato prosystemin genomic DNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       O3-JAN-2002.
(BPIG-) EPIGENOMICS AG.
(CPIG-) Match
5.4%;

Query Match
5.4%;
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(EPIG-) EPIGENOMICS AG.
                                                                                                                                                                                                                                                                             03-JAN-2002.
(EPIG-) EPIGENOMICS AG.
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Best Local Similarity
RESULT 469
Best Local Similarity
RESULT 463
                                                                                                                                                                                                                                                                                                               Best Local Similarity RESULT 466
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Best Local Similarity
RESULT 470
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                                                                                           Query Match
Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                        Local Similarity
                                                     WO9902690-A1.
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Human immune/haematopoietic antigen genomic sequence SEQ ID NO:23028.
WO200157182-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             5.4%; Score 38.6; DB 10; Length 22118; 59.6%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 22118;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 22118;
                                                                                                                                                                                                                                                                                       5.4%; Score 38.6; DB 8; Length 22118; 59.6%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                          DB 9; Length 22118;
                                                                                                       Length 21537;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           90-201.
(HUMA-) HUMAN GENOME SCI INC.
5.4%; Score 38.4; DB 4; Length 1402;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 27-SEP-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
ery Match 5.4%; Score 38.4; DB 5; Length 539;
er Thoral Similarity 53.3%; Pred. No. 32;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 637;
ABL33999 standard; DNA; 21537 BP.
Human immune system associated gene SEQ ID NO: 1972.
W0280200928-A2.
03-JAN-2002.
(EPIG) EDIGENOMICS AG.
5.44; Score 38.6; DB 6; Lensty Match
5t Local Similarity 47.3%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAL13462 standard; cDNA; 637 BP.
Human breast cancer expressed polynucleotide 5919
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB 10;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             DB 10;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             WO200151628-A2.
19-JUL-2001.
(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              (CHRO-) CHROMOS MOLECULAR SYSTEMS INC. (CHRO-) CHROMOS MOLECULAR SYSTEMS INC. 5.4%; Score 38.6; I car match 59.6%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                          5.4%; Score 38.6; I
59.6%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.4%; Score 38.6; 1
59.6%; Pred. No. 63;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       AAD61410 standard; DNA; 22118 BP.
Mouse ribosomal DNA (rDNA) repeat region.
US2003101480-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 ADL43972 standard; DNA; 539 BP.
Human ovarian cancer DNA marker #17862.
WO200170979-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Mouse ribosomal RNA gene repeat unit. W02003903469-A2. NOV-2003 (CHRO-) CHROMOS MOLECULAR SYSTEMS INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DNA; 22118 BP.
                                                                                                                                                                                                                                                                                                                                                       ADA37415 standard; DNA; 22118 BP.
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ID ADF10518 standard; DNA; 22118 BP.
                                                                                                                                                                 ADA14746 standard; DNA; 22118 BP.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                59.6%;
                                                                                                                                                                                                                                                                                                                                                                         Origin of replication DNA.
US2003108914-A1.
12-JUN-2003.
(HADL/) HADLACZKY G.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            C9-MAY-2-
(HADL/) HADLACZKY G.
(SZAL/) SZALAY A A.
                                                                                                                                                                                                                             31-OCT-2002.
(HADL/) HADLACZKY G.
(SZAL/) SZALAY A A.
                                                                                                                                                                                    Mouse 458 rRNA gene.
US2002160970-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Best Local Similarity
RESULT 475
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Best Local Similarity
RESULT 478
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Best Local Similarity
RESULT 473
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Best Local Similarity
RESULT 477
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RESULT 480
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us-09-989-293a-376.rng.spdi

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Query Match
Best Local Si
RESULT 489
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                                               (GETH )
                                                                                                           5.4%; Score 38.4; DB 3; Length 1743; 52.5%; Pred. No. 41;
                                                                                                                                                                                                                                   5.4%; Score 38.4; DB 4; Length 1743; 52.5%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                              DB 4; Length 1743;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Human angiogenesis related cDNA PRO1337 SEQ ID NO: 223 WO200208284-A2.
                                          Human PRO1337 (UNQ692) CDNA sequence SEQ 1D NO:235-WO200012708-A2.
                                                                                                                                                AAF54367 standard; DNA; 1743 BP.
Primer #140 used in the identification of proteins.
WO200078961-A1.
28-DEC-2000.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                            Human DNA encoding PRO polypeptide sequence #226.
WO200168848-A2.
20-SEP-2001.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  DB 6;
                                                                                                                                                                                                                                                                                                                                                          52.5%; Score 38.4; I 52.5%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Score 38.4; I
Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                5.4%; Score 38.4;
52.5%; Pred. No. 41;
   44.3%; Pred. No. 39;
                                                                                                                                                                                                                                                                                                                                                                                                        ABL88183 standard; cDNA; 1743 BP.
Human PRO1337 cDNA sequence SEQ ID NO:223.
WO200200690-A2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ACA89600 standard; cDNA; 1743 BP.
cDNA encoding human PRO polypeptide #226.
US2003036141-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                ABK11752 standard; cDNA; 1743 BP. cDNA encoding human PRO1337 protein. WO200149715-A2.
                             AAA37089 standard; cDNA; 1743 BP
                                                                                                                                                                                                                                                                               AAS46150 standard; cDNA; 1743 BP.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        52.58;
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(BAKE) BAKER K P.
(FERR) FERRARA N.
(GERR) GERBER H.
(GEND) GODDARD A.
(GODD) GODDARD A.
(GODO) GODOWSKI P J.
(HILL) HILLAN K J.
(HILL) HILLAN K J.
(MARS) MARSTERS S A.
(PANJ) PAN J.
                                                                          09-MAR-2000.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(STEP/) STEPHAN J F.
(WATA/) WATANABE C K.
(WILL/) WILLIAMS P M.
                                                                                                                                                                                                                                                                                                                                            (GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Best Local Similarity
RESULT 488
                                                                                                                      Best Local Similarity RESULT 482
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Best Local Similarity
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Best Local Similarity
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                                                                                                                                                                                                                                                                                                                                                                                                                                                         03-JAN-2002
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Best Local S
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              RESULT 481
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003040064-A1.
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                                                                                                                                                                                                                                                                                                                                                                                                         ACF20334 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
US2003040063-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACF00325 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
US2003054474-A1.
                                                                           Length 1743;
                                                                                                                                                                                                                 DB 8; Length 1743;
                                                                                                                                                                                                                                                                                                                                                             DB 8; Length 1743;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DB 8; Length 1743;
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US2003032114-A1.
Human secreted/transmembrane protein (PRO) cDNA #226.
US2003036146-A1.
20-FEB-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Human secreted/transmembrane protein (PRO) cDNA #226.
US2003044925-A1,
06-MAR-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Human secreted/transmembrane protein (PRO) cDNA #226.
US2003027267-A1.
                                                                                                                         ACA05925 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA
US2003036162-A1.
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                                                                         DB
                                                                                                                                                                                                               5.4%; Score 38.4; I 52.5%; Pred. No. 41;
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                                                                         Score 38.4; I
Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   52.5%; Score 38.4; I
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
                                                                                                                                                                                                                                                                                  cDNA encoding human PRO protein #226.
US2003036137-A1.
20-FEB-2003.
                                                                                                                                                                                                                                                                   CDNA; 1743 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACF19720 standard; cDNA; 1743 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ACD22008 standard; cDNA; 1743 BP
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                                                                       5.4%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         52.5%;
                                                                                                                                                                           (GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                       GENENTECH INC.
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RESULT 497
ID ACA72382 standard; cl
DE Novel human secreted
PN US200303214-A1.
                                                                                                                                                                                                                                                                                                                                                                      Best Local Similarity RESULT 491
                                                                                                                                                                                                                                  Best Local Similarity
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RESULT 495
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Best Local Similarity
RESULT 496
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Best Local Similarity
RESULT 498
                                                                                        Local Similarity
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Best Local Similarity
RESULT 494
                                                                                                                                                                                                                                                                   ACA66759 standard;
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US2003036148-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
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RESULT 504
ID ACC74387 standard; cDNA; 1743 BP.
DE Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
PN US2003027275-A1.
PD 06-FEB-2003.
5.4%; Score 38.4; DB 8; Length 1743;
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                                                                                                                                                                                                                                                                            5.4%; Score 38.4; DB 8; Length 1743; 52.5%; Pred. No. 41;
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Novel human secreted and transmembrane protein PRO1337 cDNA.
105200302294-A1.
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ID ACD25583 standard; cDNA; 1743 BP.

DE Novel human secreted and transmembrane protein PRO1337 cDNA
ACD04906 standard; cDNA; 1743 BP.
Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003032101-A1.
13-FEB-2003.
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US2003027324-A1.
06-FEB-2003.
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                                                                                                         ACD18367 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226.
20-FEB-2003.
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Human secreted/transmembrane protein (PRO) cDNA #226.
27-FEB-2003.
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Pred. No. 41;
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                                                           5.4%; Score 38.4; DB 8; 52.5%; Pred. No. 41;
                                                                                                                                                                     Score 38.4; DB 8;
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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ACA88808 standard; cDNA; 1743 BP
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ID ACD16015 standard; cDNA; 1743 BP
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20-FEB-2003.
(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 507
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Best Local Similarity
RESULT 502
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Best Local Similarity
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Best Local S
RESULT 499
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RESULT 500
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451.
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                                                                                                                                                                                                                             Human cDNA encoding a secreted/transmembrane protein, SEQ ID US2003036156-A1.
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                DB 8; Length 1743;
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Human cDNA encoding secreted/transmembrane protein, PRO1337
10520030222298-Al.
30-JAN-2003.
                                                                               Human secreted/transmembrane protein (PRO) cDNA #226. US2003040060-A1. 27-FEB-2003.
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Human secreted/transmembrane protein (PRO) cDNA #226
20.00303128-A1.
20-FEB-2003.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003027266-A1.
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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                52.5%; Score 38.4; I 52.5%; Pred. No. 41;
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ID ACD14129 standard; cDNA; 1743 BP. DE Human PRO polynucleotide #226.
PN US2003032117-A1.
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RESULT 514
ID ACD21394 standard; CDNA; 1743 BP.
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                                                                           ACD21701 standard; cDNA; 1743 BP.
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(GETH ) GENENTECH INC.
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US2003036143-A1.
20-FEB-2003.
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RESULT 512
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Best Local Similarity
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RESULT 516
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RESULT 517
ID ACA97445 standard;
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                                 Best Local Similarity RESULT 509
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20-FEB-2003.
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ACC86317 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027263-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027271-A1.
       DB 8; Length 1743;
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2002032108-A1.
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US2003036125-A1.
20-FEB-2003.
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       52.5%; Score 38.4; I 52.5%; Pred. No. 41;
                                                                                                                               Score 38.4; IPred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
                                              Human PRO polynucleotide #226.
US200302295-Al.
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(GETH ) GENENTECH INC.
 Query Match
Best Local Similarity
RESULT 529
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RESULT 535
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Best Local Similarity
RESULT 536
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RESULT 530
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RESULT 531
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RESULT 534
ID ACA70864 standard;
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Best Local Similarity
                                                                                                                                                                                   ACA89207 standard;
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RESULT 537
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003032135-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027268-A1.
06-FEB-2003.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003027274-A1.
                                                                                                                                                                             ACC91219 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003032138-A1.
                                                                                                                                                                                                                                                                                                                      ACCE8961 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003036132-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003032137-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
     5.4%; Score 38.4; DB 8; Length 1743; 52.5%; Pred. No. 41;
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                                                                                                                               DB 8; Length 1743;
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                                                                                                                                                                                                                                                                          DB 8;
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52.5%; Pred. No. 41;
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                                                     ACD14436 standard; cDNA; 1743 BP. Muman PRO polymucleotide #226. US2003032130-A1.
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Human PRO polynucleotide #226.
US2003008353-A1.
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Human PRO polynucleotide #226.
US2003017542-A1.
23-JAN-2003.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 521
Query Match
Best Local Similarity
RESULT 519
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RESULT 524
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RESULT 527
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RESULT 525
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RESULT 526
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Best Local Similarity
RESULT 520
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06-FEB-2003.
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Length 1743;

Length 1743

Length 1743;

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US2003036155-A1.
            20-FEB-2003,
           ACF20027 standard; cDNA; 1743 BP. Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.0S2003040068-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
13-2003032120-A1.
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                                                                                                                                                       Score 38.4; DB 8; Length 1743; Pred. No. 41;
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Novel human secreted and transmembrane protein PRO1337 cDNA.
                                                                                                                                                                                       ACA73303 standard; cDNA; 1743 BP.

Novel human secreted and transmembrane protein PRO1337 cDNA.
W US2003022200-A1.
D 30-JAN-2003.
S-4%; Score 38.4; DB 8; Length 1743
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003036136-Al.
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US2003332109-A1.
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
                                                               Score 38.4; DB 8;
Pred. No. 41;
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US2003036138-A1.
20-FEB-2003.
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                                                                                                ABX76971 standard; cDNA; 1743 BP.
Human PRO polynucleotide #226.
US2003027280-A1.
06-FEB-2003.
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RESULT 540
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  RESULT 539
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003032139-A1.
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                                                                                                                                                                                                                                                                                                                SEQ ID NO:451
5.4%; Score 38.4; DB 8; Length 1743; 52.5%; Pred. No. 41;
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ACA01 human secreted and transmembrane protein PRO1337 cDNA.
US2003032131-A1.
                                                                 CDNA
                                                ACD04599 standard; cDNA; 1743 BP.
Novel human secreted and transmembrane protein PRO1337
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                                                                                                                                                                                                                                                                                              ACF12702 standard; cDNA; 1743 BP.
human secreted polypeptide PRO1337-encoding cDNA,
US2003040058-A1.
27-FRB-2003.
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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CDNA encoding human PRO polypeptide #226.
US2003032107-A1.
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US2003017540-Al.
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Human PRO polynucleotide #226.
US2003032106-A1.
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             Best Local Similarity
RESULT 550
ID ACD04599 standard;
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RESULT 551
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RESULT 553
ID ACA96417 standard;
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Best Local Similarity
RESULT 556
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Best Local Similarity
RESULT 558
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RESULT 559
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                                                                                 US2003022296-A1.
30-JAN-2003.
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RESULT 571
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003027262-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003040069-A1.
27-FEB-2003.
                                                                                                                                                                                                                                                                                                                               ACC87426 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003036165-A1.
Score 38.4; DB 8; Length 1743;
Pred. No. 41;
                                                                                                                                                                                                                                                           Length 1743;
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003036154-A1.
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Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003036153-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003036142-A1.
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                                                                                                                                                                                                                                                       52.5%; Score 38.4; 1 52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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                                                                                           ACD02861 standard; cDNA; 1743 BP.
CDNA encoding human PRO polypeptide #226.
30-JAN-2003.
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13-EFF A. 13-EF
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Human PRO polynucleotide #226.
US2003036145-A1.
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52.5%;
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(GETH ) GENENTECH INC.
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                          Best Local Similarity
RESULT 560
                                                                                                                                                                                                                                                                                 Best Local Similarity
RESULT 561
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RESULT 563
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RESULT 566
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RESULT 569
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RESULT 567
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Best Local Similarity
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ACC90912 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003032122-A1.
                                                                  Length 1743;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  мылыбыл standard; cDNA; 1743 BP.
Human cDNA encoding secreted/transmembrane protein #226.
US2002127584-Al.
                                                                                                                                                                                             Human secreted/transmembrane protein (PRO) cDNA #226. US2003022116-Al.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       RESULT 576

ID ACD16629 standard; cDNA; 1743 BP.

BE Human secreted/transmembrane protein (PRO) cDNA #226.

PN US203017543 Al.

PD 23-JAN-2003.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003036152-A1.
                                                               DB 8;
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                                                                                                                                                                        Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
                                                                Score 38.4; | Pred. No. 41;
                                                                                                                                                                                                                                                                                   Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
ACA74997 standard; cDNA; 1743 BP.

CDNA encoding human PRO polypeptide #226.

1 US2003022293-A1.

0 30-JAN-2003.

5.4%; Score 38.4;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CDNA encoding human PRO polypeptide #226. US2003017541-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                          cDNA encoding human PRO protein #226. US2003036139-Al. 20-FEB-2003.
                                                                                                          ACA91868 standard; cDNA; 1743 BP.
Human PRO polynucleotide #226.
W22003032128-Al.
13-FEB-2003.
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US2003032115-A1.
13-FEB-2003.
                                                                5.4%;
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                Best Local Similarity RESULT 573
                                                                             Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                         Best Local Similarity
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                                                                                                                                                                                       Best Local Similarity
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RESULT 580
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ID ABX16811 standard;
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52.5%; Pred. No. 41;

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RESULT 581.

ID ACC1838 standard; cDNA; 1743 BP.

DE Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.

PN US200304076-A1.

PD 27-FEB-2003.

PA (GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ACF16550 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003054455-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003049741-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003049743-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ACC95987 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003036135-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
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                                             Length 1743;
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                                                                                                                                                                    Length 1743;
Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003032140-A1.
13-FBB-2003.
                                                                                                                                                                                                                  ACD11244 standard; cDNA; 1743 BP.
Novel human secreted and transmembrane protein PRO1337 cDNA.
US2003008352-A1.
                                                                                                                                                                                                                                                                                                                                     ACD15094 standard; cDNA; 1743 BP.

Human secreted/transmembrane protein (PRO) cDNA #226.
US2003044922-A1.
US-00303.
                                                                                                                                                                                                                                                                                                                                                                                                                                            ACDI1858 standard; cDNA; 1743 BP.
Human secreted/transmembrane protein (PRO) cDNA #226.
120503332118-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 38.4; DB 9;
Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              5.4%; Score 38.4; DB 9; 52.5%; Pred. No. 41;
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Pred. No. 41;
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                                                 DB 9;
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                                                                                                                                                                                                                                                                                         Score 38.4; DB 9;
Pred. No. 41;
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Pred. No. 41;
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                                                                                                                                                                      5.4%; Score 38.4; 1 52.5%; Pred. No. 41;
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                                                 Score 38.4;
Pred. No. 41;
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                                                 52.5%;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                              09-JAN-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                               Best Local Similarity
RESULT 582
ID ACD11244 standard; CD
DE Novel human secreted
PN US2003008352-A1.
PD 09-UAN-2003.
PA (GETH ) GENENTECH INC
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RESULT 589
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Best Local
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ACF78139 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US20054479-A1.
                                    ACF10246 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068743-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQ ID NO:451
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                                                                                                                     Length 1743;
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                                                                                                                                                                                                                                                             Length 1743;
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US2003068725-A1.
10-APR-2003.
(GETH ) GENENTECH INC.
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US2003068685-A1.
10-APR-2003.
(GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003068752-A1.
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                                                                                                                      Score 38.4; DB 9;
Pred. No. 41;
                                                                                                                                                                                                                                                             DB 9;
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Pred. No. 41;
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Best Local Similarity 52.5%; Pred. No. 41;
RESULT 595
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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US2003068701-A1.
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                                                                                                                        52.5%;
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Best Local Similarity 52.5%;
RESULT 597
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 591
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RESULT 593
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Best Local Similarity
RESULT 590
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RESULT 592
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Length 1743;

DB 9;

Length 1743;

DB 9;

Length 1743;

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Length 1743;

DB 9;

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ACF52045 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003064440-A1.
                                                                                                                                                                                                                                                                                Automotes Standard; cDNA; 1743 BP. Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.10-APR-2003.
                                                                                                                                                                       ACF26532 standard; cDNa; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068704-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003073183-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003064462-Al.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003049749-A1.
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                                                                                                                              38.4; D
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(GETH ) GENENTECH INC.
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... F2 5%; Pred. No. 41;
                Score 38.4; I
Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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03-5xPR-2003.
(GETH ) GENENTECH INC.
5.4%; Score 38.4;
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Pred.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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                Query Match
Best Local Similarity
RESULT 609
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RESULT 615
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Best Local Similarity
RESULT 617
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003104539-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003054473-A1.
                                                                                                                                                                                                                      ACF12088 standard; cDNA; 1743 BP.

Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
                                                                                                                                                                                                                                                                                                                                                ACF41322 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
US2003054459-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
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Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
US2003068705-A1.
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                                                                                                                                         Human secreted/transmembrane protein (PRO) cDNA #226 US2003036131-A1.
                                                                                                                                                                                        Score 38.4; DB 9;
Pred. No. 41;
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Pred. No. 41;
                                                                            Score 38.4; DB 9;
Pred. No. 41;
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Pred. No. 41;
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Human secreted/transmembrane protein (PRO) cDNA
UJS200305471-A1.
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
                                                                                                                          CDNA; 1743 BP
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(GETH ) GENENTECH INC.
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RESULT 605
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RESULT 606
                                                                                       Best_Local Similarity
RESULT 600
                                                                                                                         ACD09295 standard;
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Best Local Similarity
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RESULT 601
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                                                          05-JUN-2003.
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NO:451

Length 1743;

DB 9;

Length 1743

DB 9;

Length 1743;

DB 9;

SEQ ID NO:451

Length 1743;

DB 9;

DB 9; Length 1743;

DB 9; Length 1743;

RESULT 607

RESULT 608

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BEE
                                     Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003073184-A1.
17-APR-2003.
(GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068707-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003073175-A1.
                                                                                                                                         ACF40131 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO9741-encoding cDNA, SEQ ID NO:525.
US2003064463-A1.
                                                                                                                                                                                                                            Best Local Similarity 52.5%; Pred. No. 41;
RESULT 620
ID ACT40094 standard, cDNA, 1743 BP.
DE Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
PN US2003064463-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068721-A1.
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                                                                                                 Length 1743;
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003064451-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003068695-A1.
                                                                                                Score 38.4; DB 9;
Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               5.4%; Score 38.4; DB 9; 52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
                           ACF23404 standard; cDNA; 1743 BP
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 619
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RESULT 623
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Best Local Similarity
RESULT 625
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RESULT 626
Best Local Similarity
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Best Local Similarity
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Best Local Similarity
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PD 17
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ACC94903 standard, cDNA, 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003054468-A1.
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                                                                                                                                                                                                                                                                                 ACF77218 standard; cDNA; 1743 BP.
Human secreted polypeptide PR01337-encoding cDNA, SEQ ID NO:451
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Human secreted/transmembrane protein (PRO) cDNA #226.
20.FEB-2003.
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                                                                                                                                                                                                                                                                                                                                                                                            Best Local Similarity 52.5%; Pred. No. 41;
RESULT 630

ID ACF76911 standard; cDNA; 1743 BP.
DE Human secreted polypeptide PR01337-encoding cDNA,
PN US2003104548-A1.
PD 05-JUN-2003.
PA (GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003036130-A1.
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                                                                                             Score 38.4; DB 9;
Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
                                                                                                                                                                     cDNA encoding human PRO polypeptide #226 US2003068732-A1.
ACD84766 standard; cDNA; 1743 BP. Human PRO polymucleotide #226. US2003068703.Al. 10-APR-2003. (GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 628
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Best Local Similarity
RESULT 629
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RESULT 636
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Length 1743;

DB 9;

Length 1743;

DB 9;

Best Loca RESULT 639

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ACF25611 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068727-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003069407-Al.
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 Human secreted/transmembrane protein (PRO) cDNA #226.
US2003064461-A1.
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                                                                                                                                                                                                                                                                                                                                                                                                       ACF29295 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, US20039872-A1.
10.APR-2003.
(GETH ) GENENTECH INC.
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Pred. No. 41;
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Pred. No. 41;
                                                                                                                                                                                                              52.5%; Score 38.4; 1 52.5%; Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
                                                                                                                         ACD67803 standard; cDNA; 1743 BP.
cDNA encoding human PRO polypeptide #226.
US2003068724-A1.
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US2003068758-A1.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 654
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RESULT 647
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RESULT 651
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RESULT 655
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                                                                                                                                                                                                                                                                                   ACC97417 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003044929-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ACF14094 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003064465-A1.
                                                                                                                                                           ACF15322 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003044917-A1.
                                                                                                                                                                                                                                                                                                                                                                                                         ACC92447 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     ACF14401 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003054478-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068718-A1.
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                                                                                                                         Score 38.4; DB 9; Length 1743;
Pred. No. 41;
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Novel human secreted and transmembrane protein PRO1337 cDNA
US2003073130-A1.
                                                    Human secreted/transmembrane protein (PRO) cDNA #226.
US2003054470-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226
US2003064454-A1.
                                                                                                                                                                                                                                               Score 38.4; DB 9;
Pred. No. 41;
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Pred. No. 41;
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No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            5.4%; Score 38.4; 52.5%; Pred. No. 41;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             52.5%; Score 38.4; 52.5%; Pred. No. 41;
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Pred. No. 41;
     41;
   52.5%; Pred. No.
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                                      ACD22622 standard; cDNA; 1743 BP
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(GETH ) GENENTECH INC.
                                                                                       20-MAR-2003.
(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 646
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Best Local Similarity
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Best Local Similarity RESULT 637
                                                                                                                       Query Match
Best Local Similarity
RESULT 638
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Best Local Similarity
RESULT 640
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(GETH ) GENENTECH
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DB 9; Length 1743;

DB 9; Length 1743;

Length 1743;

DB 9;

Best Loca RESULT 644

DB 9; Length 1743;

DB 9; Length 1743;

SEQ ID NO:451

Length 1743

DB 9;

DB 9; Length 1743;

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(GETH ) GENENTECH INC.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003068709-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                       ACF08711 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
US2003049778-A1.
13-MAR-2003.
(GETH ) GENENTECH INC.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ACF38924 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068692-A1.
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Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003054476-A1.
                                                                                                                                                                                                                                                                                                                                           Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451 US2003054481-A1.
                                                                                                                                                                    Score 38.4; DB 9; Length 1743;
Pred. No. 41;
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                                                DB 9; Length 1743;
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US2003032134-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.

1052003068733-A1.

1042PR-2003.

4 (GFH) GENENTECH INC.

5.4%; Score 38.4; DB 9; Leng
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Pred. No. 41;
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Pred. No. 41;
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                                               5.4%; Score 38.4; I
52.5%; Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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Pred. No. 41;
                                                                                                             cDNA encoding human PRO polypeptide #226. US2003049771-A1.
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RESULT 658
ID ACF18264 standard; cDNA; 1743 BP.
                                                                                              ACH12400 standard; cDNA; 1743 BP.
                                                                                                                                                                                                       RESULT 657
ID ACD40792 standard; cDNA; 1743 BP
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ACF31512 standard; cDNA; 1743 BP
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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RESULT 661
ID ACF52352 standard, CDN
DE Human secreted polyper
PD 20-MAR-2003.
PA (GETH ) GENENTECH INC.
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                                  (GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
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                                                              Best Local Similarity
 US2003049768-A1.
13-MAR-2003.
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RESULT 660
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Query Match
Best Local Similarity 52.5%; Pred. No. 41;
RESULT 665
DE ACE74939 standard; cDNA; 1743 BP.
DE Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451.
PN US2003068716-A1.
                                                                                                                                                                                                                                                                                                                                                 ACF28067 standard; cDNA; 1743 BP.
Human secreted polypeptide PRO1337-encoding cDNA, SEQ ID NO:451
US2003068751-A1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ACF60583 standard; cDNA; 1743 BP. Human secreted polypeptide PR01337-encoding cDNA; SEQ ID NO:451 US2003087374-A1. 08-MAY-2003.
                                                                                                                                                                                                                     SEQ ID NO:451
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US2003073179-A1.
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Human secreted/transmembrane protein (PRO) cDNA #226.
US2003040055-A1.
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Human secreted polypeptide PRO1337-encoding cDNA,
US2003068740-A1.
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                                                                                                                                               DB 9;
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52.5%; Pred. No. 41;
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Pred. No. 41;
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Pred. No. 41;
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52.5%; Pred. No. 41;
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2.5%; Pred. No. 41;
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Pred. No. 41;
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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(GETH ) GENENTECH INC.
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Best Local Similarity
RESULT 672
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Best Local Similarity
RESULT 667
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Best Local Similarity
RESULT 670
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Best Local Similarity
RESULT 671
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RESULT 673
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